Zelig v4.0-10 Core Model Reference Manual

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1 gamma: Gamma Regression for Continuous, Positive Dependent Variables

Use the gamma regression model if you have a positive-valued dependent variable such as the number of years a parliamentary cabinet endures, or the seconds you can stay airborne while jumping. The gamma distribution assumes that all waiting times are complete by the end of the study (censoring is not allowed).

1.1 Syntax

```
> z.out <- zelig(Y ~ X1 + X2, model = "gamma", data = mydata)
> x.out <- setx(z.out)
> s.out <- sim(z.out, x = x.out, x1 = NULL)</pre>
```

1.2 Additional Inputs

In addition to the standard inputs, zelig() takes the following additional options for gamma regression:

• robust: defaults to FALSE. If TRUE is selected, zelig() computes robust standard errors via the sandwich package (see [16]). The default type of robust standard error is heteroskedastic and autocorrelation consistent (HAC), and assumes that observations are ordered by time index.

In addition, robust may be a list with the following options:

- method: Choose from
 - * "vcovHAC": (default if robust = TRUE) HAC standard errors.
 - * "kernHAC": HAC standard errors using the weights given in [1].
 - * "weave": HAC standard errors using the weights given in [11].
- order.by: defaults to NULL (the observations are chronologically ordered as in the original data). Optionally, you may specify a vector of weights (either as order.by = z, where z exists outside the data frame; or as order.by = ~z, where z is a variable in the data frame). The observations are chronologically ordered by the size of z.

- ...: additional options passed to the functions specified in method. See the sandwich library and [16] for more options.

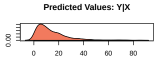
1.3 Example

```
Attach the sample data:
> data(coalition)
Estimate the model:
> z.out <- zelig(duration ~ fract + numst2, model = "gamma", data = coalition)
The following object(s) are masked from 'package:boot':
   polar
How to cite this model in Zelig:
 Kosuke Imai, Gary King, and Olivia Lau. 2012.
  "gamma: Gamma Regression for Continuous, Positive Dependent Variables"
 in Kosuke Imai, Gary King, and Olivia Lau, "Zelig: Everyone's Statistical Software,"
 http://gking.harvard.edu/zelig
View the regression output:
> summary(z.out)
Call:
glm(formula = formula, family = Gamma(), data = data, model = F)
Deviance Residuals:
   Min 1Q Median
                               3Q
                                       Max
-2.2510 -0.9112 -0.2278 0.4132
                                    1.5360
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) -1.296e-02 1.329e-02 -0.975 0.33016
            1.149e-04 1.723e-05 6.668 1.19e-10 ***
fract
numst2
           -1.739e-02 5.881e-03 -2.957 0.00335 **
Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1
(Dispersion parameter for Gamma family taken to be 0.6291004)
   Null deviance: 300.74 on 313 degrees of freedom
Residual deviance: 272.19 on 311 degrees of freedom
AIC: 2428.1
```

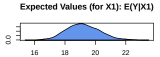
Number of Fisher Scoring iterations: 6

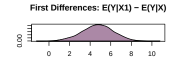
Set the baseline values (with the ruling coalition in the minority) and the alternative values (with the ruling coalition in the majority) for X:

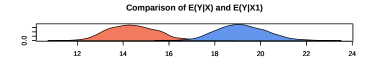
```
> x.low <- setx(z.out, numst2 = 0)
> x.high <- setx(z.out, numst2 = 1)</pre>
Simulate expected values (qi$ev) and first differences (qi$fd):
> s.out < sim(z.out, x = x.low, x1 = x.high)
> summary(s.out)
Model: gamma
Number of simulations: 1000
Values of X
  (Intercept) fract numst2
  1 718.8121 0
attr(,"assign")
[1] 0 1 2
Values of X1
  (Intercept)
             fract numst2
         1 718.8121 1
attr(,"assign")
[1] 0 1 2
Expected Values: E(Y|X)
        sd 50% 2.5% 97.5%
14.456 1.077 14.396 12.596 16.752
Expected Values (for X1): E(Y|X1)
        sd 50% 2.5% 97.5%
19.182 1.058 19.123 17.319 21.341
Predicted Values: Y|X
  mean sd 50% 2.5% 97.5%
13.889 12.24 10.315 0.646 45.575
Predicted Values: Y|X1
        sd
                50% 2.5% 97.5%
19.857 17.393 15.114 1.061 62.234
First Differences: E(Y|X1) - E(Y|X)
 mean sd 50% 2.5% 97.5%
4.725 1.497 4.78 1.745 7.524
> plot(s.out)
```











1.4 Model

• The Gamma distribution with scale parameter α has a $stochastic\ component$:

$$\begin{array}{lcl} Y & \sim & \operatorname{Gamma}(y_i \mid \lambda_i, \alpha) \\ f(y) & = & \frac{1}{\alpha^{\lambda_i} \, \Gamma \lambda_i} \, y_i^{\lambda_i - 1} \exp - \left\{ \frac{y_i}{\alpha} \right\} \end{array}$$

for $\alpha, \lambda_i, y_i > 0$.

ullet The $systematic\ component$ is given by

$$\lambda_i = \frac{1}{x_i \beta}$$

1.5 Quantities of Interest

• The expected values (qi\$ev) are simulations of the mean of the stochastic component given draws of α and β from their posteriors:

$$E(Y) = \alpha \lambda_i$$
.

- The predicted values (qi\$pr) are draws from the gamma distribution for each given set of parameters (α, λ_i) .
- If x1 is specified, sim() also returns the differences in the expected values (qi\$fd),

$$E(Y \mid x_1) - E(Y \mid x)$$

.

• In conditional prediction models, the average expected treatment effect (att.ev) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - E[Y_i(t_i=0)] \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $E[Y_i(t_i = 0)]$, the counterfactual expected value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

• In conditional prediction models, the average predicted treatment effect (att.pr) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - Y_i(\widehat{t_i=0}) \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $Y_i(\widehat{t_i} = 0)$, the counterfactual predicted value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

1.6 Output Values

The output of each Zelig command contains useful information which you may view. For example, if you run z.out <- zelig(y ~ x, model = "gamma", data), then you may examine the available information in z.out by using names(z.out), see the coefficients by using z.out\$coefficients, and a default summary of information through summary(z.out). Other elements available through the \$ operator are listed below.

- From the zelig() output object z.out, you may extract:
 - $\boldsymbol{-}$ $\mathtt{coefficients}:$ parameter estimates for the explanatory variables.

- residuals: the working residuals in the final iteration of the IWLS fit.
- fitted.values: the vector of fitted values.
- linear.predictors: the vector of $x_i\beta$.
- aic: Akaike's Information Criterion (minus twice the maximized loglikelihood plus twice the number of coefficients).
- df.residual: the residual degrees of freedom.
- df.null: the residual degrees of freedom for the null model.
- zelig.data: the input data frame if save.data = TRUE.
- From summary(z.out), you may extract:
 - coefficients: the parameter estimates with their associated standard errors, p-values, and t-statistics.
 - cov.scaled: a $k \times k$ matrix of scaled covariances.
 - cov.unscaled: a $k \times k$ matrix of unscaled covariances.
- From the sim() output object s.out, you may extract quantities of interest arranged as matrices indexed by simulation × x-observation (for more than one x-observation). Available quantities are:
 - qi\$ev: the simulated expected values for the specified values of x.
 - qi\$pr: the simulated predicted values drawn from a distribution defined by (α, λ_i) .
 - qifd: the simulated first difference in the expected values for the specified values in x and x1.
 - qi\$att.ev: the simulated average expected treatment effect for the treated from conditional prediction models.
 - qi\$att.pr: the simulated average predicted treatment effect for the treated from conditional prediction models.

How to Cite the Gamma Model

How to Cite the Zelig Software Package

To cite Zelig as a whole, please reference these two sources:

Kosuke Imai, Gary King, and Olivia Lau. 2007. "Zelig: Everyone's Statistical Software," http://GKing.harvard.edu/zelig.

Imai, Kosuke, Gary King, and Olivia Lau. (2008). "Toward A Common Framework for Statistical Analysis and Development." Journal of Computational and Graphical Statistics, Vol. 17, No. 4 (December), pp. 892-913.

See also

The gamma model is part of the stats package by [14]. Advanced users may wish to refer to help(glm) and help(family), as well as [13]. Robust standard errors are implemented via the sandwich package by [16]. Sample data are from [9].

2 logit: Logistic Regression for Dichotomous Dependent Variables

Logistic regression specifies a dichotomous dependent variable as a function of a set of explanatory variables.

2.1 Syntax

```
> z.out <- zelig(Y ~ X1 + X2, model = "logit", data = mydata)
> x.out <- setx(z.out)
> s.out <- sim(z.out, x = x.out, x1 = NULL)</pre>
```

2.2 Additional Inputs

In addition to the standard inputs, zelig() takes the following additional options for logistic regression:

• robust: defaults to FALSE. If TRUE is selected, zelig() computes robust standard errors via the sandwich package (see [16]). The default type of robust standard error is heteroskedastic and autocorrelation consistent (HAC), and assumes that observations are ordered by time index.

In addition, robust may be a list with the following options:

- method: Choose from
 - * "vcovHAC": (default if robust = TRUE) HAC standard errors.
 - * "kernHAC": HAC standard errors using the weights given in [1].
 - * "weave": HAC standard errors using the weights given in [11].
- order.by: defaults to NULL (the observations are chronologically ordered as in the original data). Optionally, you may specify a vector of weights (either as order.by = z, where z exists outside the data frame; or as order.by = ~z, where z is a variable in the data frame)
 The observations are chronologically ordered by the size of z.
- ...: additional options passed to the functions specified in method.
 See the sandwich library and [16] for more options.

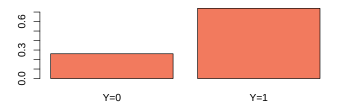
2.3 Examples

1. Basic Example Attaching the sample turnout dataset: > data(turnout) Estimating parameter values for the logistic regression: > z.out1 <- zelig(vote ~ age + race, model = "logit", data = turnout) How to cite this model in Zelig: Kosuke Imai, Gary King, and Olivia Lau. 2012. "logit: Logistic Regression for Dichotomous Dependent Variables" in Kosuke Imai, Gary King, and Olivia Lau, "Zelig: Everyone's Statistical Software," http://gking.harvard.edu/zelig Setting values for the explanatory variables: > x.out1 <- setx(z.out1, age = 36, race = "white")</pre> Simulating quantities of interest from the posterior distribution. > s.out1 <- sim(z.out1, x = x.out1) > summary(s.out1) Model: logit Number of simulations: 1000 Values of X (Intercept) age racewhite 1 36 attr(,"assign") [1] 0 1 2 attr(,"contrasts") attr(,"contrasts")\$race [1] "contr.treatment" Expected Values: E(Y|X)sd 50% 2.5% 97.5% 0.748 0.012 0.748 0.724 0.77 Predicted Values: Y|X

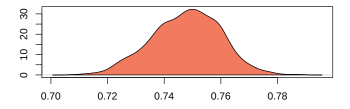
0 1 0.261 0.739

> plot(s.out1)

Predicted Values: Y|X



Expected Values: E(Y|X)



2. Simulating First Differences

Estimating the risk difference (and risk ratio) between low education (25th percentile) and high education (75th percentile) while all the other variables held at their default values.

```
> z.out2 <- zelig(vote ~ race + educate, model = "logit", data = turnout)
```

How to cite this model in Zelig:

Kosuke Imai, Gary King, and Olivia Lau. 2012.

"logit: Logistic Regression for Dichotomous Dependent Variables" in Kosuke Imai, Gary King, and Olivia Lau, "Zelig: Everyone's Statistical Software,"

In Kosuke Imai, Gary King, and Ulivia Lau, "Zelig: Everyone's Statistical Software http://gking.harvard.edu/zelig

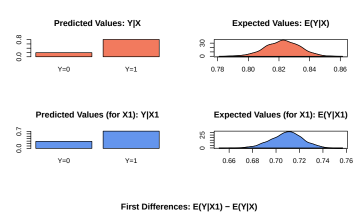
- > x.high <- setx(z.out2, educate = quantile(turnout\$educate, prob = 0.75))
- > x.low <- setx(z.out2, educate = quantile(turnout\$educate, prob = 0.25))</pre>
- > s.out2 <- sim(z.out2, x = x.high, x1 = x.low)
- > summary(s.out2)

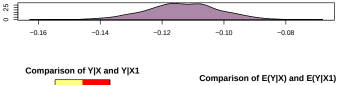
Model: logit Number of simulations: 1000 Values of X (Intercept) racewhite educate 1 1 1 attr(,"assign") [1] 0 1 2 attr(,"contrasts") attr(,"contrasts")\$race [1] "contr.treatment" Values of X1 (Intercept) racewhite educate 1 10 attr(,"assign") [1] 0 1 2 attr(,"contrasts") attr(,"contrasts")\$race [1] "contr.treatment" Expected Values: E(Y|X)sd 50% 2.5% 97.5% mean 0.822 0.011 0.823 0.8 0.843 Expected Values (for X1): E(Y|X1) sd 50% 2.5% 97.5% 0.709 0.013 0.71 0.682 0.734 Predicted Values: Y|X 0 1 0.191 0.809 Predicted Values (for X1): Y|X1 0 0.289 0.711 First Differences: E(Y|X1) - E(Y|X)

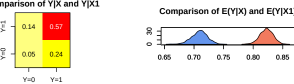
> plot(s.out2)

sd 50% 2.5% 97.5%

-0.113 0.012 -0.113 -0.137 -0.091







3. Presenting Results: An ROC Plot

One can use an ROC plot to evaluate the fit of alternative model specifications. (Use demo(roc) to view this example, or see King and Zeng (2002).)

```
> z.out1 <- zelig(vote ~ race + educate + age, model = "logit",
+ data = turnout)</pre>
```

How to cite this model in Zelig:

Kosuke Imai, Gary King, and Olivia Lau. 2012.

"logit: Logistic Regression for Dichotomous Dependent Variables"

in Kosuke Imai, Gary King, and Olivia Lau, "Zelig: Everyone's Statistical Software," http://gking.harvard.edu/zelig

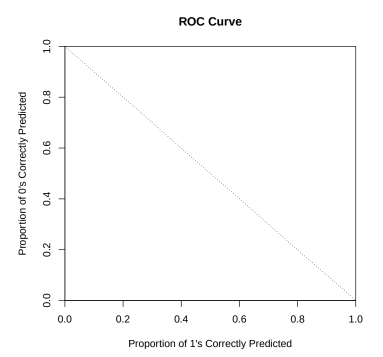
> z.out2 <- zelig(vote ~ race + educate, model = "logit", data = turnout)

How to cite this model in Zelig:

Kosuke Imai, Gary King, and Olivia Lau. 2012.

"logit: Logistic Regression for Dichotomous Dependent Variables"

in Kosuke Imai, Gary King, and Olivia Lau, "Zelig: Everyone's Statistical Software," http://gking.harvard.edu/zelig



2.4 Model

Let Y_i be the binary dependent variable for observation i which takes the value of either 0 or 1.

• The stochastic component is given by

$$Y_i \sim \text{Bernoulli}(y_i \mid \pi_i)$$

= $\pi_i^{y_i} (1 - \pi_i)^{1 - y_i}$

where $\pi_i = \Pr(Y_i = 1)$.

 \bullet The *systematic component* is given by:

$$\pi_i = \frac{1}{1 + \exp(-x_i \beta)}.$$

where x_i is the vector of k explanatory variables for observation i and β is the vector of coefficients.

2.5 Quantities of Interest

• The expected values (qi\$ev) for the logit model are simulations of the predicted probability of a success:

$$E(Y) = \pi_i = \frac{1}{1 + \exp(-x_i \beta)},$$

given draws of β from its sampling distribution.

- The predicted values (qi\$pr) are draws from the Binomial distribution with mean equal to the simulated expected value π_i .
- The first difference (qi\$fd) for the logit model is defined as

$$FD = Pr(Y = 1 \mid x_1) - Pr(Y = 1 \mid x).$$

• The risk ratio (qi\$rr) is defined as

$$RR = Pr(Y = 1 \mid x_1) / Pr(Y = 1 \mid x).$$

• In conditional prediction models, the average expected treatment effect (att.ev) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - E[Y_i(t_i=0)] \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $E[Y_i(t_i = 0)]$, the counterfactual expected value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

• In conditional prediction models, the average predicted treatment effect (att.pr) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - \widehat{Y_i(t_i=0)} \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $Y_i(\widehat{t_i} = 0)$, the counterfactual predicted value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

2.6 Output Values

The output of each Zelig command contains useful information which you may view. For example, if you run z.out <- zelig(y ~ x, model = "logit", data), then you may examine the available information in z.out by using names(z.out), see the coefficients by using z.out\$coefficients, and a default summary of information through summary(z.out). Other elements available through the \$ operator are listed below.

- From the zelig() output object z.out, you may extract:
 - coefficients: parameter estimates for the explanatory variables.
 - residuals: the working residuals in the final iteration of the IWLS fit.
 - fitted.values: the vector of fitted values for the systemic component, π_i .
 - linear.predictors: the vector of $x_i\beta$
 - aic: Akaike's Information Criterion (minus twice the maximized loglikelihood plus twice the number of coefficients).
 - df.residual: the residual degrees of freedom.
 - df.null: the residual degrees of freedom for the null model.
 - data: the name of the input data frame.
- From summary(z.out), you may extract:
 - coefficients: the parameter estimates with their associated standard errors, p-values, and t-statistics.
 - cov.scaled: a $k \times k$ matrix of scaled covariances.
 - cov.unscaled: a $k \times k$ matrix of unscaled covariances.
- From the sim() output object s.out, you may extract quantities of interest arranged as matrices indexed by simulation × x-observation (for more than one x-observation). Available quantities are:
 - qi\$ev: the simulated expected probabilities for the specified values of x.
 - qi\$pr: the simulated predicted values for the specified values of x.
 - qi\$fd: the simulated first difference in the expected probabilities for the values specified in x and x1.
 - qi\$rr: the simulated risk ratio for the expected probabilities simulated from x and x1.
 - qi\$att.ev: the simulated average expected treatment effect for the treated from conditional prediction models.
 - qi\$att.pr: the simulated average predicted treatment effect for the treated from conditional prediction models.

3 ls: Least Squares Regression for Continuous Dependent Variables

Use least squares regression analysis to estimate the best linear predictor for the specified dependent variables.

3.1 Syntax

```
> z.out <- zelig(Y ~ X1 + X2, model = "ls", data = mydata)
> x.out <- setx(z.out)
> s.out <- sim(z.out, x = x.out)</pre>
```

3.2 Additional Inputs

In addition to the standard inputs, zelig() takes the following additional options for least squares regression:

• robust: defaults to FALSE. If TRUE is selected, zelig() computes robust standard errors based on sandwich estimators (see [16], [2], and [15]). The default type of robust standard error is heteroskedastic consistent (HC), not heteroskedastic and autocorrelation consistent (HAC).

In addition, robust may be a list with the following options:

- method: choose from
 - * "vcovHC": (the default if robust = TRUE), HC standard errors.
 - * "vcovHAC": HAC standard errors without weights.
 - * "kernHAC": HAC standard errors using the weights given in [1].
 - * "weave": HAC standard errors using the weights given in [11].
- order.by: only applies to the HAC methods above. Defaults to NULL (the observations are chronologically ordered as in the original data).
 Optionally, you may specify a time index (either as order.by = z, where z exists outside the data frame; or as order.by = ~z, where z is a variable in the data frame). The observations are chronologically ordered by the size of z.
- ...: additional options passed to the functions specified in method.
 See the sandwich library and [16] for more options.

3.3 Examples

1. Basic Example with First Differences

Attach sample data:

> data(macro)

Estimate model:

```
How to cite this model in Zelig:
  Kosuke Imai, Gary King, and Olivia Lau. 2012.
  "ls: Least Squares Regression for Continuous Dependent Variables"
  in Kosuke Imai, Gary King, and Olivia Lau, "Zelig: Everyone's Statistical Software,"
  http://gking.harvard.edu/zelig
Summarize regression coefficients:
> summary(z.out1)
Call:
lm(formula = formula, data = data, weights = weights, model = F)
Residuals:
   Min
             1Q Median
                             3Q
                                     Max
-5.3008 -2.0768 -0.3187 1.9789 7.7715
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) 6.181294 0.450572 13.719 < 2e-16 ***
                       0.062820 -5.151 4.36e-07 ***
gdp
            -0.323601
capmob
             1.421939
                       0.166443
                                   8.543 4.22e-16 ***
             0.019854 0.005606
                                   3.542 0.000452 ***
trade
Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1
Residual standard error: 2.746 on 346 degrees of freedom
Multiple R-squared: 0.2878,
                                   Adjusted R-squared: 0.2817
F-statistic: 46.61 on 3 and 346 DF, p-value: < 2.2e-16
Set explanatory variables to their default (mean/mode) values, with high
(80th percentile) and low (20th percentile) values for the trade variable:
> x.high <- setx(z.out1, trade = quantile(macro$trade, 0.8))
> x.low <- setx(z.out1, trade = quantile(macro$trade, 0.2))
Generate first differences for the effect of high versus low trade on GDP:
> s.out1 <- sim(z.out1, x = x.high, x1 = x.low)
> summary(s.out1)
Model: ls
Number of simulations: 1000
Values of X
```

> z.out1 <- zelig(unem ~ gdp + capmob + trade, model = "ls", data = macro)

```
(Intercept)
                           capmob
                                     trade
                   gdp
            1 3.254223 -0.8914286 79.10131
1
attr(,"assign")
[1] 0 1 2 3
Values of X1
  (Intercept)
                           capmob
                                     trade
                   gdp
            1 3.254223 -0.8914286 37.29106
attr(,"assign")
[1] 0 1 2 3
Expected Values: E(Y|X)
   mean
          sd 50% 2.5% 97.5%
1 5.439 0.204 5.434 5.045 5.832
Expected Values (of X1): E(Y|X1)
 mean
          sd 50% 2.5% 97.5%
1 4.6 0.183 4.597 4.256 4.972
First Difference in Expected Values: E(Y|X1) - E(Y|X)
                 50%
                       2.5% 97.5%
    mean
            sd
1 -0.839 0.234 -0.844 -1.294 -0.383
```

2. Using Dummy Variables

Estimate a model with fixed effects for each country. Note that you do not need to create dummy variables, as the program will automatically parse the unique values in the selected variable into discrete levels.

Set values for the explanatory variables, using the default mean/mode values, with country set to the United States and Japan, respectively:

```
> x.US <- setx(z.out2, country = "United States")
> x.Japan <- setx(z.out2, country = "Japan")</pre>
```

Simulate quantities of interest:

> s.out2 <- sim(z.out2, x = x.US, x1 = x.Japan)

3.4 Model

• The stochastic component is described by a density with mean μ_i and the common variance σ^2

$$Y_i \sim f(y_i \mid \mu_i, \sigma^2).$$

• The systematic component models the conditional mean as

$$\mu_i = x_i \beta$$

where x_i is the vector of covariates, and β is the vector of coefficients.

The least squares estimator is the best linear predictor of a dependent variable given x_i , and minimizes the sum of squared residuals, $\sum_{i=1}^{n} (Y_i - x_i\beta)^2$.

3.5 Quantities of Interest

 The expected value (qi\$ev) is the mean of simulations from the stochastic component,

$$E(Y) = x_i \beta$$
,

given a draw of β from its sampling distribution.

 In conditional prediction models, the average expected treatment effect (att.ev) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - E[Y_i(t_i=0)] \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $E[Y_i(t_i = 0)]$, the counterfactual expected value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

3.6 Output Values

The output of each Zelig command contains useful information which you may view. For example, if you run z.out <- zelig(y ~ x, model = "ls", data), then you may examine the available information in z.out by using names(z.out) see the coefficients by using z.out\$coefficients, and a default summary of information through summary(z.out). Other elements available through the \$ operator are listed below.

- From the zelig() output object z.out, you may extract:
 - coefficients: parameter estimates for the explanatory variables.
 - residuals: the working residuals in the final iteration of the IWLS fit.
 - fitted.values: fitted values.
 - df.residual: the residual degrees of freedom.
 - zelig.data: the input data frame if save.data = TRUE.
- From summary(z.out), you may extract:
 - coefficients: the parameter estimates with their associated standard errors, p-values, and t-statistics.

$$\hat{\beta} = \left(\sum_{i=1}^{n} x_i' x_i\right)^{-1} \sum x_i y_i$$

sigma: the square root of the estimate variance of the random errore:

$$\hat{\sigma} = \frac{\sum (Y_i - x_i \hat{\beta})^2}{n - k}$$

- r.squared: the fraction of the variance explained by the model.

$$R^{2} = 1 - \frac{\sum (Y_{i} - x_{i}\hat{\beta})^{2}}{\sum (y_{i} - \bar{y})^{2}}$$

- adj.r.squared: the above R^2 statistic, penalizing for an increased number of explanatory variables.
- cov.unscaled: a $k \times k$ matrix of unscaled covariances.
- From the sim() output object s.out, you may extract quantities of interest arranged as matrices indexed by simulation × x-observation (for more than one x-observation). Available quantities are:
 - qi\$ev: the simulated expected values for the specified values of x.
 - qi\$fd: the simulated first differences (or differences in expected values) for the specified values of x and x1.
 - qi\$att.ev: the simulated average expected treatment effect for the treated from conditional prediction models.

How to Cite the Least Squares Model How to Cite the Zelig Software Package

To cite Zelig as a whole, please reference these two sources:

Kosuke Imai, Gary King, and Olivia Lau. 2007. "Zelig: Everyone's Statistical Software," http://GKing.harvard.edu/zelig.

Imai, Kosuke, Gary King, and Olivia Lau. (2008). "Toward A Common Framework for Statistical Analysis and Development." Journal of Computational and Graphical Statistics, Vol. 17, No. 4 (December), pp. 892-913.

See also

The least squares regression is part of the stats package by William N. Venables and Brian D. Ripley [14]. In addition, advanced users may wish to refer to help(lm) and help(lm.fit). Robust standard errors are implemented via the sandwich package by Achim Zeileis [16]. Sample data are from [9].

How to Cite the Logit Model

How to Cite the Zelig Software Package

To cite Zelig as a whole, please reference these two sources:

Kosuke Imai, Gary King, and Olivia Lau. 2007. "Zelig: Everyone's Statistical Software," http://GKing.harvard.edu/zelig.

Imai, Kosuke, Gary King, and Olivia Lau. (2008). "Toward A Common Framework for Statistical Analysis and Development." Journal of Computational and Graphical Statistics, Vol. 17, No. 4 (December), pp. 892-913.

See also

The logit model is part of the stats package by [14]. Advanced users may wish to refer to help(glm) and help(family), as well as [13]. Robust standard errors are implemented via the sandwich package by [16]. Sample data are from [9].

4 negbinom: Negative Binomial Regression for Event Count Dependent Variables

Use the negative binomial regression if you have a count of events for each observation of your dependent variable. The negative binomial model is frequently used to estimate over-dispersed event count models.

4.1 Syntax

```
> z.out <- zelig(Y ~ X1 + X2, model = "negbinom", data = mydata)
> x.out <- setx(z.out)
> s.out <- sim(z.out, x = x.out)</pre>
```

4.2 Additional Inputs

In addition to the standard inputs, zelig() takes the following additional options for negative binomial regression:

• robust: defaults to FALSE. If TRUE is selected, zelig() computes robust standard errors via the sandwich package (see [16]). The default type of robust standard error is heteroskedastic and autocorrelation consistent (HAC), and assumes that observations are ordered by time index.

In addition, robust may be a list with the following options:

- method: Choose from
 - * "vcovHAC": (default if robust = TRUE) HAC standard errors.
 - * "kernHAC": HAC standard errors using the weights given in [1].
 - * "weave": HAC standard errors using the weights given in [11].
- order.by: defaults to NULL (the observations are chronologically ordered as in the original data). Optionally, you may specify a vector of weights (either as order.by = z, where z exists outside the data frame; or as order.by = z, where z is a variable in the data frame). The observations are chronologically ordered by the size of z.
- ...: additional options passed to the functions specified in method. See the sandwich library and [16] for more options.

4.3 Example

```
Load sample data:

> data(sanction)

Estimate the model:

> z.out <- zelig(num ~ target + coop, model = "negbinom", data = sanction)

The following object(s) are masked from 'package:MASS':

coop

How to cite this model in Zelig:
```

```
Kosuke Imai, Gary King, and Olivia Lau. 2012.
"negbinom: Negative Binomial Regression for Event Count Dependent Variables"
in Kosuke Imai, Gary King, and Olivia Lau, "Zelig: Everyone's Statistical Software,"
http://gking.harvard.edu/zelig
```

```
> summary(z.out)
Call:
glm.nb(formula = num ~ target + coop, data = Data.frame, weights = NULL,
    init.theta = 1.841603403, link = log)
Deviance Residuals:
   Min
        1Q Median 3Q
                                       Max
-2.0302 -0.5118 -0.1418 -0.0191
                                    3.9987
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) -1.5641
                        0.5941 -2.633 0.0103 *
target
             0.1510
                         0.2262
                                0.668 0.5063
             1.2857
                        0.1761 7.302 2.51e-10 ***
coop
Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1
(Dispersion parameter for Negative Binomial(1.8416) family taken to be 1.520566)
    Null deviance: 237.094 on 77 degrees of freedom
Residual deviance: 56.545 on 75 degrees of freedom
AIC: 360.19
Number of Fisher Scoring iterations: 1
Set values for the explanatory variables to their default mean values:
> x.out <- setx(z.out)</pre>
Simulate fitted values:
> s.out <- sim(z.out, x = x.out)
> summary(s.out)
Model: negbinom
Number of simulations: 1000
Values of X
  (Intercept) target
                          coop
           1 2.141026 1.807692
1
attr(,"assign")
[1] 0 1 2
Expected Values: E(Y|X)
```

50% 2.5% 97.5%

sd

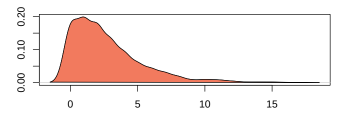
2.996 0.453 2.968 2.233 4.028

Predicted Values: Y|X

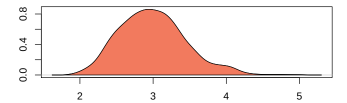
mean sd 50% 2.5% 97.5% 2.731 2.699 2 0 10

> plot(s.out)

Predicted Values: Y|X



Expected Values: E(Y|X)



4.4 Model

Let Y_i be the number of independent events that occur during a fixed time period. This variable can take any non-negative integer value.

• The negative binomial distribution is derived by letting the mean of the Poisson distribution vary according to a fixed parameter ζ given by the Gamma distribution. The *stochastic component* is given by

$$Y_i \mid \zeta_i \sim \operatorname{Poisson}(\zeta_i \mu_i),$$

 $\zeta_i \sim \frac{1}{\theta} \operatorname{Gamma}(\theta).$

The marginal distribution of Y_i is then the negative binomial with mean

 μ_i and variance $\mu_i + \mu_i^2/\theta$:

$$Y_i \sim \operatorname{NegBinom}(\mu_i, \theta),$$

$$= \frac{\Gamma(\theta + y_i)}{y! \Gamma(\theta)} \frac{\mu_i^{y_i} \theta^{\theta}}{(\mu_i + \theta)^{\theta + y_i}},$$

where θ is the systematic parameter of the Gamma distribution modeling ζ_i .

• The systematic component is given by

$$\mu_i = \exp(x_i \beta)$$

where x_i is the vector of k explanatory variables and β is the vector of coefficients.

4.5 Quantities of Interest

• The expected values (qi\$ev) are simulations of the mean of the stochastic component. Thus,

$$E(Y) = \mu_i = \exp(x_i \beta),$$

given simulations of β .

- The predicted value (qi\$pr) drawn from the distribution defined by the set of parameters (μ_i, θ) .
- The first difference (qi\$fd) is

$$FD = E(Y|x_1) - E(Y \mid x)$$

• In conditional prediction models, the average expected treatment effect (att.ev) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - E[Y_i(t_i=0)] \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $E[Y_i(t_i = 0)]$, the counterfactual expected value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

• In conditional prediction models, the average predicted treatment effect (att.pr) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - \widehat{Y_i(t_i=0)} \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $\widehat{Y_i(t_i = 0)}$, the counterfactual predicted value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

4.6 Output Values

The output of each Zelig command contains useful information which you may view. For example, if you run z.out <- zelig(y ~ x, model = "negbinom", data), then you may examine the available information in z.out by using names(z.out), see the coefficients by using z.out\$coefficients, and a default summary of information through summary(z.out). Other elements available through the \$ operator are listed below.

- From the zelig() output object z.out, you may extract:
 - coefficients: parameter estimates for the explanatory variables.
 - theta: the maximum likelihood estimate for the stochastic parameter θ .
 - SE.theta: the standard error for theta.
 - residuals: the working residuals in the final iteration of the IWLS fit
 - fitted.values: a vector of the fitted values for the systemic component λ .
 - linear.predictors: a vector of $x_i\beta$.
 - aic: Akaike's Information Criterion (minus twice the maximized loglikelihood plus twice the number of coefficients).
 - df.residual: the residual degrees of freedom.
 - df.null: the residual degrees of freedom for the null model.
 - zelig.data: the input data frame if save.data = TRUE.
- From summary(z.out), you may extract:
 - coefficients: the parameter estimates with their associated standard errors, p-values, and t-statistics.
 - cov.scaled: a $k \times k$ matrix of scaled covariances.
 - cov.unscaled: a $k \times k$ matrix of unscaled covariances.
- From the sim() output object s.out, you may extract quantities of interest arranged as matrices indexed by simulation × x-observation (for more than one x-observation). Available quantities are:

- qi\$ev: the simulated expected values given the specified values of x.
- qi\$pr: the simulated predicted values drawn from the distribution defined by (μ_i, θ) .
- qi\$fd: the simulated first differences in the simulated expected values given the specified values of x and x1.
- qi\$att.ev: the simulated average expected treatment effect for the treated from conditional prediction models.
- qi\$att.pr: the simulated average predicted treatment effect for the treated from conditional prediction models.

How to Cite the Negative Binomial Model How to Cite the Zelig Software Package

To cite Zelig as a whole, please reference these two sources:

Kosuke Imai, Gary King, and Olivia Lau. 2007. "Zelig: Everyone's Statistical Software," http://GKing.harvard.edu/zelig.

Imai, Kosuke, Gary King, and Olivia Lau. (2008). "Toward A Common Framework for Statistical Analysis and Development." Journal of Computational and Graphical Statistics, Vol. 17, No. 4 (December), pp. 892-913.

5 normal: Normal Regression for Continuous Dependent Variables

The Normal regression model is a close variant of the more standard least squares regression model (see Section 3). Both models specify a continuous dependent variable as a linear function of a set of explanatory variables. The Normal model reports maximum likelihood (rather than least squares) estimates. The two models differ only in their estimate for the stochastic parameter σ .

5.1 Syntax

```
> z.out <- zelig(Y ~ X1 + X2, model = "normal", data = mydata)
> x.out <- setx(z.out)
> s.out <- sim(z.out, x = x.out)</pre>
```

5.2 Additional Inputs

In addition to the standard inputs, zelig() takes the following additional options for normal regression:

• robust: defaults to FALSE. If TRUE is selected, zelig() computes robust standard errors via the sandwich package (see [16]). The default type of robust standard error is heteroskedastic and autocorrelation consistent (HAC), and assumes that observations are ordered by time index.

In addition, robust may be a list with the following options:

- method: Choose from
 - * "vcovHAC": (default if robust = TRUE) HAC standard errors.
 - * "kernHAC": HAC standard errors using the weights given in [1].
 - * "weave": HAC standard errors using the weights given in [11].
- order.by: defaults to NULL (the observations are chronologically ordered as in the original data). Optionally, you may specify a vector of weights (either as order.by = z, where z exists outside the data frame; or as order.by = ~z, where z is a variable in the data frame). The observations are chronologically ordered by the size of z.
- ...: additional options passed to the functions specified in method. See the sandwich library and [16] for more options.

5.3 Examples

1. Basic Example with First Differences

Attach sample data:

```
> data(macro)
```

Estimate model:

Summarize of regression coefficients:

```
> summary(z.out1)
Call:
glm(formula = formula, family = gaussian, data = data, weights = weights,
    model = F)
Deviance Residuals:
    Min    1Q    Median    3Q    Max
```

```
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) 6.181294
                         0.450572 13.719 < 2e-16 ***
                         0.062820 -5.151 4.36e-07 ***
gdp
            -0.323601
                         0.166443
                                    8.543 4.22e-16 ***
capmob
             1.421939
                                    3.542 0.000452 ***
                         0.005606
trade
             0.019854
Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1
(Dispersion parameter for gaussian family taken to be 7.54307)
    Null deviance: 3664.8 on 349
                                    degrees of freedom
Residual deviance: 2609.9 on 346 degrees of freedom
AIC: 1706.5
Number of Fisher Scoring iterations: 2
Set explanatory variables to their default (mean/mode) values, with high
(80th percentile) and low (20th percentile) values for trade:
> x.high <- setx(z.out1, trade = quantile(macro$trade, 0.8))
> x.low <- setx(z.out1, trade = quantile(macro$trade, 0.2))</pre>
Generate first differences for the effect of high versus low trade on GDP:
> s.out1 <- sim(z.out1, x = x.high, x1 = x.low)
> summary(s.out1)
Model: normal
Number of simulations: 1000
Values of X
  (Intercept)
                    gdp
                            capmob
                                      trade
            1 3.254223 -0.8914286 79.10131
attr(,"assign")
[1] 0 1 2 3
Values of X1
  (Intercept)
                            capmob
                                      trade
                   gdp
            1 3.254223 -0.8914286 37.29106
attr(,"assign")
[1] 0 1 2 3
Expected Values: E(Y|X)
```

1.9789

7.7715

-5.3008 -2.0768 -0.3187

mean sd 50% 2.5% 97.5% 5.428 0.193 5.43 5.065 5.815

Expected Values (for X1): E(Y|X1) mean sd 50% 2.5% 97.5% 4.604 0.178 4.598 4.258 4.945

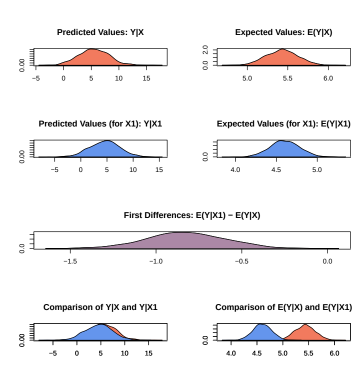
Predicted Values: Y|X mean sd 50% 2.5% 97.5% 5.452 2.699 5.473 0.197 11.161

Predicted Values (for X1): Y|X1 mean sd 50% 2.5% 97.5% 4.543 2.833 4.685 -1.012 10.221

First Differences: E(Y|X1) - E(Y|X) mean sd 50% 2.5% 97.5% -0.824 0.231 -0.826 -1.274 -0.383

A visual summary of quantities of interest:

> plot(s.out1)



2. Using Dummy Variables

Estimate a model with a dummy variable for each year and country (see factors for help with dummy variables). Note that you do not need to create dummy variables, as the program will automatically parse the unique values in the selected variables into dummy variables.

Set values for the explanatory variables, using the default mean/mode variables, with country set to the United States and Japan, respectively:

```
> ### x.US <- try(setx(z.out2, country = "United States"), silent=T)
> ### x.Japan <- try(setx(z.out2, country = "Japan"), silent=T)
Simulate quantities of interest:
> ### s.out2 <- try(sim(z.out2, x = x.US, x1 = x.Japan), silent=T)</pre>
```

> ###try(summary(s.out2))

5.4 Model

Let Y_i be the continuous dependent variable for observation i.

• The stochastic component is described by a univariate normal model with a vector of means μ_i and scalar variance σ^2 :

$$Y_i \sim \text{Normal}(\mu_i, \sigma^2).$$

• The *systematic component* is

$$\mu_i = x_i \beta$$
,

where x_i is the vector of k explanatory variables and β is the vector of coefficients.

5.5 Quantities of Interest

• The expected value (qi\$ev) is the mean of simulations from the the stochastic component,

$$E(Y) = \mu_i = x_i \beta,$$

given a draw of β from its posterior.

- The predicted value (qi\$pr) is drawn from the distribution defined by the set of parameters (μ_i, σ) .
- The first difference (qi\$fd) is:

$$FD = E(Y \mid x_1) - E(Y \mid x)$$

• In conditional prediction models, the average expected treatment effect (att.ev) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - E[Y_i(t_i=0)] \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $E[Y_i(t_i = 0)]$, the counterfactual expected value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

• In conditional prediction models, the average predicted treatment effect (att.pr) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - Y_i(\widehat{t_i=0}) \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $Y_i(\widehat{t_i} = 0)$, the counterfactual predicted value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

5.6 Output Values

The output of each Zelig command contains useful information which you may view. For example, if you run z.out <- zelig(y ~ x, model = "normal", data), then you may examine the available information in z.out by using names(z.out), see the coefficients by using z.out\$coefficients, and a default summary of information through summary(z.out). Other elements available through the \$ operator are listed below.

- From the zelig() output object z.out, you may extract:
 - coefficients: parameter estimates for the explanatory variables.
 - residuals: the working residuals in the final iteration of the IWLS fit.
 - fitted.values: fitted values. For the normal model, these are identical to the linear predictors.
 - linear.predictors: fitted values. For the normal model, these are identical to fitted.values.
 - aic: Akaike's Information Criterion (minus twice the maximized loglikelihood plus twice the number of coefficients).
 - df.residual: the residual degrees of freedom.
 - df.null: the residual degrees of freedom for the null model.
 - zelig.data: the input data frame if save.data = TRUE.
- From summary(z.out), you may extract:
 - coefficients: the parameter estimates with their associated standard errors, p-values, and t-statistics.
 - cov.scaled: a $k \times k$ matrix of scaled covariances.
 - cov.unscaled: a $k \times k$ matrix of unscaled covariances.
- From the sim() output object s.out, you may extract quantities of interest arranged as matrices indexed by simulation × x-observation (for more than one x-observation). Available quantities are:
 - qi\$ev: the simulated expected values for the specified values of x.
 - qi\$pr: the simulated predicted values drawn from the distribution defined by (μ_i, σ) .
 - qi\$fd: the simulated first difference in the simulated expected values for the values specified in x and x1.
 - qi\$att.ev: the simulated average expected treatment effect for the treated from conditional prediction models.
 - qi\$att.pr: the simulated average predicted treatment effect for the treated from conditional prediction models.

How to Cite the Normal Regression Model How to Cite the Zelig Software Package

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Imai, Kosuke, Gary King, and Olivia Lau. (2008). "Toward A Common Framework for Statistical Analysis and Development." Journal of Computational and Graphical Statistics, Vol. 17, No. 4 (December), pp. 892-913.

See also

The normal model is part of the stats package by [14]. Advanced users may wish to refer to help(glm) and help(family), as well as [13]. Robust standard errors are implemented via the sandwich package by [16]. Sample data are from [9].

6 poisson: Poisson Regression for Event Count Dependent Variables

Use the Poisson regression model if the observations of your dependent variable represents the number of independent events that occur during a fixed period of time (see the negative binomial model, Section 4, for over-dispersed event counts.).

6.1 Syntax

```
> z.out <- zelig(Y ~ X1 + X2, model = "poisson", data = mydata)
> x.out <- setx(z.out)
> s.out <- sim(z.out, x = x.out)</pre>
```

6.2 Additional Inputs

In addition to the standard inputs, zelig() takes the following additional options for poisson regression:

• robust: defaults to FALSE. If TRUE is selected, zelig() computes robust standard errors via the sandwich package (see [16]). The default type of robust standard error is heteroskedastic and autocorrelation consistent (HAC), and assumes that observations are ordered by time index.

In addition, robust may be a list with the following options:

- method: Choose from
 - * "vcovHAC": (default if robust = TRUE) HAC standard errors.
 - * "kernHAC": HAC standard errors using the weights given in [1].
 - * "weave": HAC standard errors using the weights given in [11].
- order.by: defaults to NULL (the observations are chronologically ordered as in the original data). Optionally, you may specify a vector of weights (either as order.by = z, where z exists outside the data

```
frame; or as order.by = ~z, where z is a variable in the data frame).
The observations are chronologically ordered by the size of z.
```

- ...: additional options passed to the functions specified in method. See the sandwich library and [16] for more options.

6.3

```
Example
Load sample data:
> data(sanction)
Estimate Poisson model:
> z.out <- zelig(num ~ target + coop, model = "poisson", data = sanction)
The following object(s) are masked from 'package:MASS':
    coop
 How to cite this model in Zelig:
 Kosuke Imai, Gary King, and Olivia Lau. 2012.
  "poisson: Poisson Regression for Event Count Dependent Variables"
  in Kosuke Imai, Gary King, and Olivia Lau, "Zelig: Everyone's Statistical Software,"
 http://gking.harvard.edu/zelig
> summary(z.out)
Call:
glm(formula = formula, family = poisson(), data = data, weights = weights,
   model = F)
Deviance Residuals:
         1Q Median
                               3Q
                                       Max
-7.2127 -1.1831 -0.2080 -0.1856 17.6514
Coefficients:
           Estimate Std. Error z value Pr(>|z|)
(Intercept) -0.96772
                       0.17545 -5.516 3.48e-08 ***
                       0.05823 -0.361
target
           -0.02102
                                          0.718
            1.21082
                       0.04662 25.970 < 2e-16 ***
coop
Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1
(Dispersion parameter for poisson family taken to be 1)
```

Null deviance: 1583.77 on 77 degrees of freedom

```
Residual deviance: 720.84 on 75 degrees of freedom
```

AIC: 944.35

Number of Fisher Scoring iterations: 6

Set values for the explanatory variables to their default mean values:

> x.out <- setx(z.out)</pre>

Simulate fitted values:

> s.out <- sim(z.out, x = x.out)

> summary(s.out)

Model: poisson

Number of simulations: 1000

Values of X

(Intercept) target coop 1 1 2.141026 1.807692

attr(,"assign")

[1] 0 1 2

Expected Values: E(Y|X)

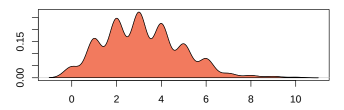
mean sd 50% 2.5% 97.5% 3.248 0.234 3.245 2.798 3.745

Predicted Values: Y|X

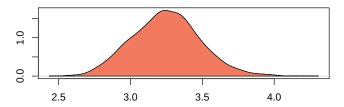
mean sd 50% 2.5% 97.5% 3.172 1.738 3 0 7

> plot(s.out)

Predicted Values: Y|X



Expected Values: E(Y|X)



6.4 Model

Let Y_i be the number of independent events that occur during a fixed time period. This variable can take any non-negative integer.

• The Poisson distribution has stochastic component

$$Y_i \sim \text{Poisson}(\lambda_i),$$

where λ_i is the mean and variance parameter.

• The systematic component is

$$\lambda_i = \exp(x_i \beta),$$

where x_i is the vector of explanatory variables, and β is the vector of coefficients.

6.5 Quantities of Interest

• The expected value (qi\$ev) is the mean of simulations from the stochastic component,

$$E(Y) = \lambda_i = \exp(x_i \beta),$$

given draws of β from its sampling distribution.

- The predicted value (qi\$pr) is a random draw from the poisson distribution defined by mean λ_i .
- The first difference in the expected values (qi\$fd) is given by:

$$FD = E(Y|x_1) - E(Y \mid x)$$

• In conditional prediction models, the average expected treatment effect (att.ev) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - E[Y_i(t_i=0)] \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $E[Y_i(t_i = 0)]$, the counterfactual expected value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

• In conditional prediction models, the average predicted treatment effect (att.pr) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t=1}^{n} \left\{ Y_i(t_i = 1) - Y_i(\widehat{t_i} = 0) \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $Y_i(\widehat{t_i} = 0)$, the counterfactual predicted value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

6.6 Output Values

The output of each Zelig command contains useful information which you may view. For example, if you run z.out <- zelig(y ~ x, model = "poisson", data), then you may examine the available information in z.out by using names(z.out), see the coefficients by using z.out\$coefficients, and a default summary of information through summary(z.out). Other elements available through the \$ operator are listed below.

- From the zelig() output object z.out, you may extract:
 - coefficients: parameter estimates for the explanatory variables.
 - residuals: the working residuals in the final iteration of the IWLS fit.

- fitted.values: a vector of the fitted values for the systemic component λ .
- linear.predictors: a vector of $x_i\beta$.
- aic: Akaike's Information Criterion (minus twice the maximized loglikelihood plus twice the number of coefficients).
- df.residual: the residual degrees of freedom.
- df.null: the residual degrees of freedom for the null model.
- zelig.data: the input data frame if save.data = TRUE.
- From summary(z.out), you may extract:
 - coefficients: the parameter estimates with their associated standard errors, p-values, and t-statistics.
 - cov.scaled: a $k \times k$ matrix of scaled covariances.
 - cov.unscaled: a $k \times k$ matrix of unscaled covariances.
- From the sim() output object s.out, you may extract quantities of interest arranged as matrices indexed by simulation × x-observation (for more than one x-observation). Available quantities are:
 - qi\$ev: the simulated expected values given the specified values of x.
 - qi\$pr: the simulated predicted values drawn from the distributions defined by λ_i .
 - qifd: the simulated first differences in the expected values given the specified values of x and x1.
 - qi\$att.ev: the simulated average expected treatment effect for the treated from conditional prediction models.
 - qi\$att.pr: the simulated average predicted treatment effect for the treated from conditional prediction models.

How to Cite the Poisson Regression Model How to Cite the Zelig Software Package

To cite Zelig as a whole, please reference these two sources:

Kosuke Imai, Gary King, and Olivia Lau. 2007. "Zelig: Everyone's Statistical Software," http://GKing.harvard.edu/zelig.

Imai, Kosuke, Gary King, and Olivia Lau. (2008). "Toward A Common Framework for Statistical Analysis and Development." Journal of Computational and Graphical Statistics, Vol. 17, No. 4 (December), pp. 892-913.

See also

The poisson model is part of the stats package by [14]. Advanced users may wish to refer to help(glm) and help(family), as well as [13]. Robust standard errors are implemented via the sandwich package by [16]. Sample data are from [12].

7 probit: Probit Regression for Dichotomous Dependent Variables

Use probit regression to model binary dependent variables specified as a function of a set of explanatory variables.

7.1 Syntax

```
> z.out <- zelig(Y ~ X1 + X2, model = "probit", data = mydata)
> x.out <- setx(z.out)
> s.out <- sim(z.out, x = x.out, x1 = NULL)</pre>
```

7.2 Additional Inputs

In addition to the standard inputs, zelig() takes the following additional options for probit regression:

• robust: defaults to FALSE. If TRUE is selected, zelig() computes robust standard errors via the sandwich package (see [16]). The default type of robust standard error is heteroskedastic and autocorrelation consistent (HAC), and assumes that observations are ordered by time index.

In addition, robust may be a list with the following options:

- method: Choose from
 - * "vcovHAC": (default if robust = TRUE) HAC standard errors.
 - * "kernHAC": HAC standard errors using the weights given in [1].
 - * "weave": HAC standard errors using the weights given in [11].
- order.by: defaults to NULL (the observations are chronologically ordered as in the original data). Optionally, you may specify a vector of weights (either as order.by = z, where z exists outside the data frame; or as order.by = z, where z is a variable in the data frame). The observations are chronologically ordered by the size of z.
- ...: additional options passed to the functions specified in method. See the sandwich library and [16] for more options.

7.3Examples

Attach the sample turnout dataset:

```
> data(turnout)
Estimate parameter values for the probit regression:
> z.out <- zelig(vote ~ race + educate, model = "probit", data = turnout)
How to cite this model in Zelig:
 Kosuke Imai, Gary King, and Olivia Lau. 2012.
  "probit: Probit Regression for Dichotomous Dependent Variables"
  in Kosuke Imai, Gary King, and Olivia Lau, "Zelig: Everyone's Statistical Software,"
 http://gking.harvard.edu/zelig
> summary(z.out)
Call:
glm(formula = formula, family = binomial(link = "probit"), data = data,
    weights = weights, model = F)
Deviance Residuals:
   Min 10 Median
                                3Q
                                       Max
-2.2586 -0.8982 0.6712 0.7232
                                     1.7045
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) -0.725949
                       0.128635 -5.643 1.67e-08 ***
racewhite
            0.299076
                       0.084648 3.533 0.000411 ***
                       0.009571 10.147 < 2e-16 ***
educate
            0.097119
Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 2266.7 on 1999 degrees of freedom
Residual deviance: 2136.0 on 1997 degrees of freedom
AIC: 2142
Number of Fisher Scoring iterations: 4
Set values for the explanatory variables to their default values.
> x.out <- setx(z.out)
```

Simulate quantities of interest from the posterior distribution.

> s.out <- sim(z.out, x = x.out)

```
> summary(s.out)
Model: probit
Number of simulations: 1000
Values of X
  (Intercept) racewhite educate
            1
                     1 12.06675
attr(,"assign")
[1] 0 1 2
attr(,"contrasts")
attr(,"contrasts")$race
[1] "contr.treatment"
Expected Values: E(Y|X)
        sd 50% 2.5% 97.5%
 mean
 0.772 0.01 0.772 0.751 0.791
```

0.219 0.781

Model

0

7.4

Predicted Values: Y|X

Let Y_i be the observed binary dependent variable for observation i which takes the value of either 0 or 1.

• The stochastic component is given by

$$Y_i \sim \text{Bernoulli}(\pi_i),$$

where $\pi_i = \Pr(Y_i = 1)$.

• The systematic component is

$$\pi_i = \Phi(x_i\beta)$$

where $\Phi(\mu)$ is the cumulative distribution function of the Normal distribution with mean 0 and unit variance.

7.5 Quantities of Interest

 \bullet The expected value (qi\$ev) is a simulation of predicted probability of success

$$E(Y) = \pi_i = \Phi(x_i \beta),$$

given a draw of β from its sampling distribution.

- The predicted value (qi\$pr) is a draw from a Bernoulli distribution with mean π_i .
- The first difference (qi\$fd) in expected values is defined as

$$FD = Pr(Y = 1 \mid x_1) - Pr(Y = 1 \mid x).$$

• The risk ratio (qi\$rr) is defined as

$$RR = Pr(Y = 1 \mid x_1) / Pr(Y = 1 \mid x).$$

• In conditional prediction models, the average expected treatment effect (att.ev) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t_i=1}^{n} \left\{ Y_i(t_i=1) - E[Y_i(t_i=0)] \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $E[Y_i(t_i = 0)]$, the counterfactual expected value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

• In conditional prediction models, the average predicted treatment effect (att.pr) for the treatment group is

$$\frac{1}{\sum_{i=1}^{n} t_i} \sum_{i:t=1}^{n} \left\{ Y_i(t_i = 1) - \widehat{Y_i(t_i = 0)} \right\},\,$$

where t_i is a binary explanatory variable defining the treatment $(t_i = 1)$ and control $(t_i = 0)$ groups. Variation in the simulations are due to uncertainty in simulating $Y_i(\widehat{t_i} = 0)$, the counterfactual predicted value of Y_i for observations in the treatment group, under the assumption that everything stays the same except that the treatment indicator is switched to $t_i = 0$.

7.6 Output Values

The output of each Zelig command contains useful information which you may view. For example, if you run z.out <- zelig(y ~ x, model = "probit", data), then you may examine the available information in z.out by using names(z.out), see the coefficients by using z.out\$coefficients, and a default summary of information through summary(z.out). Other elements available through the \$ operator are listed below.

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- residuals: the working residuals in the final iteration of the IWLS fit.
- fitted.values: a vector of the in-sample fitted values.
- linear.predictors: a vector of $x_i\beta$.
- aic: Akaike's Information Criterion (minus twice the maximized loglikelihood plus twice the number of coefficients).
- df.residual: the residual degrees of freedom.
- df.null: the residual degrees of freedom for the null model.
- data: the name of the input data frame.
- From summary(z.out), you may extract:
 - coefficients: the parameter estimates with their associated standard errors, p-values, and t-statistics.
 - cov.scaled: a $k \times k$ matrix of scaled covariances.
 - cov.unscaled: a $k \times k$ matrix of unscaled covariances.
- From the sim() output object s.out, you may extract quantities of interest arranged as matrices indexed by simulation × x-observation (for more than one x-observation). Available quantities are:
 - qi\$ev: the simulated expected values, or predicted probabilities, for the specified values of x.
 - qi\$pr: the simulated predicted values drawn from the distributions defined by the predicted probabilities.
 - qifd: the simulated first differences in the predicted probabilities for the values specified in x and x1.
 - qi\$rr: the simulated risk ratio for the predicted probabilities simulated from x and x1.
 - qi\$att.ev: the simulated average expected treatment effect for the treated from conditional prediction models.
 - qi\$att.pr: the simulated average predicted treatment effect for the treated from conditional prediction models.

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See also

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8 Bibliography

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