0.1 poisson.survey: Survey-Weighted Poisson Regression for Event Count Dependent Variables

The survey-weighted poisson regression model is appropriate for survey data obtained using complex sampling techniques, such as stratified random or cluster sampling (e.g., not simple random sampling). Like the conventional poisson regression model (see Section ??), survey-weighted poisson regression specifies a dependent variable representing the number of independent events that occur during a fixed period of time as function of a set of explanatory variables. The survey-weighted poisson model reports estimates of model parameters identical to conventional poisson estimates, but uses information from the survey design to correct variance estimates.

The poisson.survey model accommodates three common types of complex survey data. Each method listed here requires selecting specific options which are detailed in the "Additional Inputs" section below.

- 1. **Survey weights**: Survey data are often published along with weights for each observation. For example, if a survey intentionally over-samples a particular type of case, weights can be used to correct for the over-representation of that type of case in the dataset. Survey weights come in two forms:
 - (a) *Probability* weights report the probability that each case is drawn from the population. For each stratum or cluster, this is computed as the number of observations in the sample drawn from that group divided by the number of observations in the population in the group.
 - (b) Sampling weights are the inverse of the probability weights.
- 2. **Strata/cluster identification**: A complex survey dataset may include variables that identify the strata or cluster from which observations are drawn. For stratified random sampling designs, observations may be nested in different strata. There are two ways to employ these identifiers:
 - (a) Use *finite population corrections* to specify the total number of cases in the stratum or cluster from which each observation was drawn.
 - (b) For stratified random sampling designs, use the raw strata ids to compute sampling weights from the data.
- 3. **Replication weights**: To preserve the anonymity of survey participants, some surveys exclude strata and cluster ids from the public data and instead release only precomputed replicate weights.

Syntax

```
> z.out <- zelig(Y ~ X1 + X2, model = "poisson.survey", data = mydata)
```

```
> x.out <- setx(z.out)
> s.out <- sim(z.out, x = x.out)</pre>
```

Additional Inputs

In addition to the standard zelig inputs (see Section ??), survey-weighted poisson models accept the following optional inputs:

- 1. Datasets that include survey weights:.
 - probs: An optional formula or numerical vector specifying each case's probability weight, the probability that the case was selected. Probability weights need not (and, in most cases, will not) sum to one. Cases with lower probability weights are weighted more heavily in the computation of model coefficients.
 - weights: An optional numerical vector specifying each case's sample weight, the inverse of the probability that the case was selected. Sampling weights need not (and, in most cases, will not) sum to one. Cases with higher sampling weights are weighted more heavily in the computation of model coefficients.
- 2. Datasets that include strata/cluster identifiers:
 - ids: An optional formula or numerical vector identifying the cluster from which each observation was drawn (ordered from largest level to smallest level). For survey designs that do not involve cluster sampling, ids defaults to NULL.
 - fpc: An optional numerical vector identifying each case's frequency weight, the total number of units in the population from which each observation was sampled.
 - strata: An optional formula or vector identifying the stratum from which each observation was sampled. Entries may be numerical, logical, or strings. For survey designs that do not involve cluster sampling, strata defaults to NULL.
 - nest: An optional logical value specifying whether primary sampling unites (PSUs) have non-unique ids across multiple strata. nest=TRUE is appropriate when PSUs reuse the same identifiers across strata. Otherwise, nest defaults to FALSE.
 - check.strata: An optional input specifying whether to check that clusters are nested in strata. If check.strata is left at its default, !nest, the check is not performed. If check.strata is specified as TRUE, the check is carried out.
- 3. Datasets that include replication weights:
 - repweights: A formula or matrix specifying replication weights, numerical vectors of weights used in a process in which the sample is repeatedly re-weighted and parameters are re-estimated in order to compute the variance of the model parameters.

- type: A string specifying the type of replication weights being used. This input is required if replicate weights are specified. The following types of replication weights are recognized: "BRR", "Fay", "JK1", "JKn", "bootstrap", or "other".
- weights: An optional vector or formula specifying each case's sample weight, the inverse of the probability that the case was selected. If a survey includes both sampling weights and replicate weights separately for the same survey, both should be included in the model specification. In these cases, sampling weights are used to correct potential biases in in the computation of coefficients and replication weights are used to compute the variance of coefficient estimates.
- combined.weights: An optional logical value that should be specified as TRUE if the replicate weights include the sampling weights. Otherwise, combined.weights defaults to FALSE.
- rho: An optional numerical value specifying a shrinkage factor for replicate weights of type "Fay".
- bootstrap.average: An optional numerical input specifying the number of iterations over which replicate weights of type "bootstrap" were averaged. This input should be left as NULL for "bootstrap" weights that were not were created by averaging.
- scale: When replicate weights are included, the variance is computed as the sum of squared deviations of the replicates from their mean. scale is an optional overall multiplier for the standard deviations.
- rscale: Like scale, rscale specifies an optional vector of replicate-specific multipliers for the squared deviations used in variance computation.
- fpc: For models in which "JK1", "JKn", or "other" replicates are specified, fpc is an optional numerical vector (with one entry for each replicate) designating the replicates' finite population corrections.
- fpctype: When a finite population correction is included as an fpc input, fpctype is a required input specifying whether the input to fpc is a sampling fraction (fpctype="fraction") or a direct correction (fpctype="correction").
- return.replicates: An optional logical value specifying whether the replicates should be returned as a component of the output. return.replicates defaults to FALSE.

Examples

1. A dataset that includes survey weights:

Attach the sample data:

> data(api, package = "survey")

Suppose that a dataset included a variable reporting the number of times a new student enrolled during the previous school year (enroll), a measure of each school's academic performance (api99), an indicator for whether each school holds classes year round (year.rnd), and sampling weights computed by the survey house (pw). Estimate a model that regresses enroll on api99 and year.rnd:

```
> z.out1 <- zelig(enroll ~ api99 + yr.rnd, model = "poisson.survey",
+ weights = ~pw, data = apistrat)</pre>
```

Summarize regression coefficients:

```
> summary(z.out1)
```

Set explanatory variables to their default (mean/mode) values, and set a high (80th percentile) and low (20th percentile) value for the measure of academic performance:

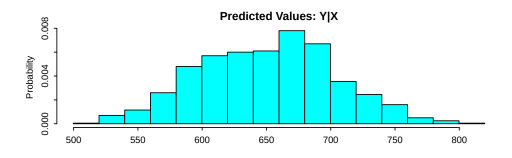
```
> x.low <- setx(z.out1, api99 = quantile(apistrat$api99, 0.2))
> x.high <- setx(z.out1, api99 = quantile(apistrat$api99, 0.8))</pre>
```

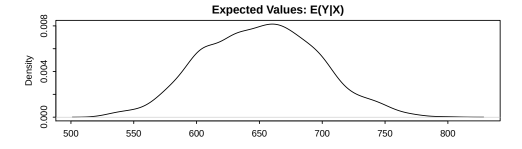
Generate first differences for the effect of high versus low concentrations of children receiving subsidized meals on the probability of holding school year-round:

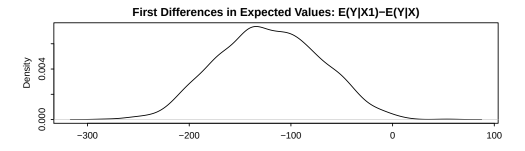
```
> s.out1 <- sim(z.out1, x = x.low, x1 = x.high)
> summary(s.out1)
```

Generate a visual summary of the quantities of interest:

```
> plot(s.out1)
```







2. A dataset that includes strata/cluster identifiers:

Suppose that the survey house that provided the dataset used in the previous example excluded sampling weights but made other details about the survey design available. A user can still estimate a model without sampling weights that instead uses inputs that identify the stratum and/or cluster to which each observation belongs and the size of the finite population from which each observation was drawn.

Continuing the example above, suppose the survey house drew at random a fixed number of elementary schools, a fixed number of middle schools, and a fixed number of high schools. If the variable stype is a vector of characters ("E" for elementary schools, "M" for middle schools, and "H" for high schools) that identifies the type of school each case represents and fpc is a numerical vector that identifies for each case the total number of schools of the same type in the population, then the user could estimate the following model:

```
> z.out2 <- zelig(enroll ~ api99 + yr.rnd, model = "poisson.survey",
+ data = apistrat, strata = ~stype, fpc = ~fpc)</pre>
```

Summarize the regression output:

```
> summary(z.out2)
```

The coefficient estimates for this example are identical to the point estimates in the first example, when pre-existing sampling weights were used. When sampling weights are omitted, they are estimated automatically for "poisson.survey" models based on the user-defined description of sampling designs.

Moreover, because the user provided information about the survey design, the standard error estimates are lower in this example than in the previous example, in which the user omitted variables pertaining to the details of the complex survey design.

3. A dataset that includes replication weights:

Consider a dataset that includes information for a sample of hospitals about the number of out-of-hospital cardiac arrests that each hospital treats and the number of patients who arrive alive at each hospital:

```
> data(scd, package = "survey")
```

Survey houses sometimes supply replicate weights (in lieu of details about the survey design). For the sake of illustrating how replicate weights can be used as inputs in normal.survey models, create a set of balanced repeated replicate (BRR) weights:

```
> BRRrep <- 2 * cbind(c(1, 0, 1, 0, 1, 0), c(1, 0, 0, 1, 0, 1),
+ c(0, 1, 1, 0, 0, 1), c(0, 1, 0, 1, 1, 0))
```

Estimate a model that regresses the count of patients who arrived alive at the hospital last year on the number of patients treated for cardiac arrests, using the BRR replicate weights in BRRrep to compute standard errors:

```
> z.out3 <- zelig(alive ~ arrests, model = "poisson.survey", repweights = BRRrep,
+ type = "BRR", data = scd)
> summary(z.out3)
```

Summarize the regression coefficients:

```
> summary(z.out3)
```

Set the explanatory variable arrests at its 20th and 80th quantiles:

```
> x.low <- setx(z.out3, arrests = quantile(scd$arrests, 0.2))
> x.high <- setx(z.out3, arrests = quantile(scd$arrests, 0.8))</pre>
```

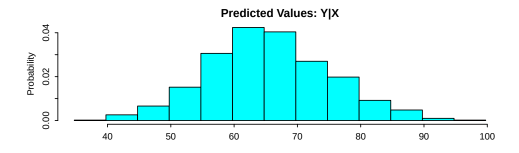
Generate first differences for the effect of high versus low cardiac arrests on the count of patients who arrive alive:

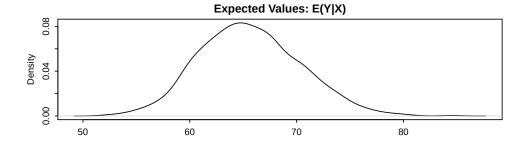
```
> s.out3 <- sim(z.out3, x = x.high, x1 = x.low)
```

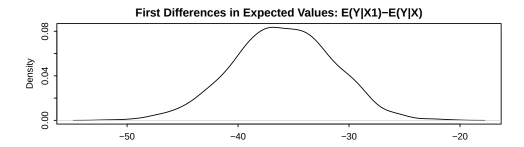
> summary(s.out3)

Generate a visual summary of quantities of interest:

> plot(s.out3)







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.

Variance

When replicate weights are not used, the variance of the coefficients is estimated as

$$\hat{\Sigma} \left[\sum_{i=1}^{n} \frac{(1-\pi_i)}{\pi_i^2} (\mathbf{X}_i(Y_i - \mu_i))' (\mathbf{X}_i(Y_i - \mu_i)) + 2 \sum_{i=1}^{n} \sum_{j=i+1}^{n} \frac{(\pi_{ij} - \pi_i \pi_j)}{\pi_i \pi_j \pi_{ij}} (\mathbf{X}_i(Y_i - \mu_i))' (\mathbf{X}_j(Y_j - \mu_j)) \right] \hat{\Sigma}$$

where π_i is the probability of case *i* being sampled, \mathbf{X}_i is a vector of the values of the explanatory variables for case *i*, Y_i is value of the dependent variable for case *i*, $\hat{\mu}_i$ is the predicted value of the dependent variable for case *i* based on the linear model estimates, and $\hat{\Sigma}$ is the conventional variance-covariance matrix in a parametric glm. This statistic is derived from the method for estimating the variance of sums described in Binder (1983) and the Horvitz-Thompson estimator of the variance of a sum described in Horvitz and Thompson (1952).

When replicate weights are used, the model is re-estimated for each set of replicate weights, and the variance of each parameter is estimated by summing the squared deviations of the replicates from their mean.

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:

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

• 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 $Y_i(t_i = 0)$, 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 $t_i = 0$.

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.survey", 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 log-likelihood 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 .
 - qi\$fd: 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.

When users estimate poisson.survey models with replicate weights in Zelig, an object called .survey.prob.weights is created in the global environment. Zelig will over-write any existing object with that name, and users are therefore advised to re-name any object called .survey.prob.weights before using poisson.survey models in Zelig.

How to Cite

To cite the *poisson.survey* Zelig model:

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See also

Survey-weighted linear models and the sample data used in the examples above are a part of the survey package by Thomas Lumley. Users may wish to refer to the help files for the three functions that Zelig draws upon when estimating survey-weighted models, namely, help(svyglm), help(svydesign), and help(svrepdesign). The Gamma model is part of the stats package by Venables and Ripley (2002). Advanced users may wish to refer to help(glm) and help(family), as well as McCullagh and Nelder (1989).

Bibliography

- Binder, D. A. (1983), "On the Variance of Asymptoticaly Normal Estimators from Complex Surveys," *International Statistical Review*, 51, 279–292.
- Horvitz, D. G. and Thompson, D. (1952), "A Generalization of Sampling without Replacement from a Finite Universe," *Journal of the American Statistical Association*, 47, 663–685.
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- Venables, W. N. and Ripley, B. D. (2002), *Modern Applied Statistics with S*, Springer-Verlag, 4th ed.