Title

tnbreg — Truncated negative binomial regression

Description Options Acknowledgment Quick start Remarks and examples References Menu Stored results Also see Syntax Methods and formulas

Description

tnbreg estimates the parameters of a truncated negative binomial model by maximum likelihood. The dependent variable *depvar* is regressed on *indepvars*, where *depvar* is a positive count variable whose values are all above the truncation point.

Quick start

Truncated negative binomial regression of y on x with truncation at 0

tnbreg y x

Report incidence-rate ratios

tnbreg y x, irr

- Add categorical variable a using factor variable syntax tnbreg y x i.a
- Same as above, but specify a constant truncation point of 2 tnbreg y x i.a, 11(2)
- With exposure variable exp tnbreg y x i.a, exposure(exp)
- Same as above, but specifying a variable truncation point stored in variable min tnbreg y x i.a, exposure(exp) ll(min)
- With cluster-robust standard errors clustering by the levels of cvar tnbreg y x i.a, exposure(exp) ll(min) vce(cluster cvar)

Menu

Statistics > Count outcomes > Truncated negative binomial regression

Syntax

tnbreg depvar [indepvars] [if] [in] [weight] [, options]

options	Description
Model	
<u>nocons</u> tant	suppress constant term
11(# <i>varname</i>)	truncation point; default value is 11(0), zero truncation
\underline{d} ispersion(\underline{m} ean)	parameterization of dispersion; the default
<u>d</u> ispersion(<u>c</u> onstant)	constant dispersion for all observations
$\underline{exposure}(varname_e)$	include $\ln(varname_e)$ in model with coefficient constrained to 1
$\underline{off}set(varname_o)$	include varname _o in model with coefficient constrained to 1
<pre><u>const</u>raints(constraints)</pre>	apply specified linear constraints
SE/Robust	
vce(vcetype)	<pre>vcetype may be oim, robust, cluster clustvar, opg, bootstrap, or jackknife</pre>
Reporting	
<u>l</u> evel(#)	set confidence level; default is level(95)
<u>nolr</u> test	suppress likelihood-ratio test
<u>ir</u> r	report incidence-rate ratios
<u>nocnsr</u> eport	do not display constraints
display_options	control columns and column formats, row spacing, line width, display of omitted variables and base and empty cells, and factor-variable labeling
Maximization	
maximize_options	control the maximization process; seldom used
<u>col</u> linear	keep collinear variables
<u>coefl</u> egend	display legend instead of statistics

indepvars may contain factor variables; see [U] 11.4.3 Factor variables.

depvar and indepvars may contain time-series operators; see [U] 11.4.4 Time-series varlists.

bayes, bootstrap, by, collect, fp, jackknife, rolling, statsby, and svy are allowed; see [U] 11.1.10 Prefix commands. For more details, see [BAYES] bayes: thbreg.

Weights are not allowed with the bootstrap prefix; see [R] bootstrap.

vce() and weights are not allowed with the svy prefix; see [SVY] svy.

fweights, iweights, and pweights are allowed; see [U] 11.1.6 weight.

collinear and coeflegend do not appear in the dialog box.

See [U] 20 Estimation and postestimation commands for more capabilities of estimation commands.

Options

Model

noconstant; see [R] Estimation options.

11(# | varname) specifies the truncation point, which is a nonnegative integer. The default is zero truncation, 11(0).

dispersion(mean | constant) specifies the parameterization of the model. dispersion(mean), the default, yields a model with dispersion equal to $1 + \alpha \exp(\mathbf{x}_j\beta + \text{offset}_j)$; that is, the dispersion is a function of the expected mean: $\exp(\mathbf{x}_j\beta + \text{offset}_j)$. dispersion(constant) has dispersion equal to $1 + \delta$; that is, it is a constant for all observations.

exposure(varname_e), offset(varname_o), constraints(constraints); see [R] Estimation options.

SE/Robust

vce(vcetype) specifies the type of standard error reported, which includes types that are derived from asymptotic theory (oim, opg), that are robust to some kinds of misspecification (robust), that allow for intragroup correlation (cluster clustvar), and that use bootstrap or jackknife methods (bootstrap, jackknife); see [R] vce_option.

Reporting

level(#); see [R] Estimation options.

- nolrtest suppresses fitting the Poisson model. Without this option, a comparison Poisson model is fit, and the likelihood is used in a likelihood-ratio test of the null hypothesis that the dispersion parameter is zero.
- irr reports estimated coefficients transformed to incidence-rate ratios, that is, e^{β_i} rather than β_i . Standard errors and confidence intervals are similarly transformed. This option affects how results are displayed, not how they are estimated or stored. irr may be specified at estimation or when replaying previously estimated results.

nocnsreport; see [R] Estimation options.

display_options: noci, nopvalues, noomitted, vsquish, noemptycells, baselevels, allbaselevels, notvlabel, fvwrap(#), fvwrapon(style), cformat(% fmt), pformat(% fmt), sformat(% fmt), and nolstretch; see [R] Estimation options.

Maximization

maximize_options: difficult, technique(algorithm_spec), iterate(#), [no]log, trace, gradient, showstep, hessian, showtolerance, tolerance(#), ltolerance(#), nrtolerance(#), nonrtolerance, and from(init_specs); see [R] Maximize. These options are seldom used.

Setting the optimization type to technique(bhhh) resets the default vcetype to vce(opg).

The following options are available with tnbreg but are not shown in the dialog box:

collinear, coeflegend; see [R] Estimation options.

Remarks and examples

Grogger and Carson (1991) showed that overdispersion causes inconsistent estimation of the mean in the truncated Poisson model. To solve this problem, they proposed using the truncated negative binomial model as an alternative. If data are truncated but do not exhibit overdispersion, the truncated Poisson model is more appropriate; see [R] **tpoisson**. For an introduction to negative binomial regression, see Cameron and Trivedi (2005, 2022) and Long and Freese (2014). For an introduction to truncated negative binomial models, see Cameron and Trivedi (2013) and Long (1997, chap. 8).

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tnbreg fits the mean-dispersion and the constant-dispersion parameterizations of truncated negative binomial models. These parameterizations extend those implemented in nbreg; see [R] nbreg.

Example 1

We illustrate the truncated negative binomial model using the 1997 MedPar dataset (Hilbe 1999). The data are from 1,495 patients in Arizona who were assigned to a diagnostic-related group (DRG) of patients having a ventilator. Length of stay (los), the dependent variable, is a positive integer; it cannot have zero values. The data are truncated because there are no observations on individuals who stayed for zero days.

The objective of this example is to determine whether the length of stay was related to the binary variables: died, hmo, type1, type2, and type3.

The died variable was recorded as a 0 unless the patient died, in which case, it was recorded as a 1. The other variables also adopted this encoding. The hmo variable was set to 1 if the patient belonged to a health maintenance organization (HMO).

The type1-type3 variables indicated the type of admission used for the patient. The type1 variable indicated an emergency admit. The type2 variable indicated an urgent admit—that is, the first available bed. The type3 variable indicated an elective admission. Because type1-type3 were mutually exclusive, only two of the three could be used in the truncated negative binomial regression shown below.

. use https:// (Arizona venti	/www.stata-pre ilator data)	ss.com/data	a/r18/medj	par		
. tnbreg los died hmo type2-type3, vce(cluster provnum) nolog						
Truncated negative binomial regression Truncation point = 0 Dispersion: mean Log pseudolikelihood = -4737.535			Number of ob Wald chi2(4) Prob > chi2 Pseudo R2	= 36.01		
		(Std. e	err. adju	sted for	54 clusters i	n provnum)
		Robust				
los	Coefficient	std. err.	Z	P> z	[95% conf.	interval]
died	2521884	.061533	-4.10	0.000	3727908	1315859
hmo	0754173	.0533132	-1.41	0.157	1799091	.0290746
type2	.2685095	.0666474	4.03	0.000	.137883	.3991359
type3	.7668101	.2183505	3.51	0.000	.338851	1.194769
_cons	2.224028	.034727	64.04	0.000	2.155964	2.292091
/lnalpha	630108	.0764019			779853	480363
alpha	.5325343	.0406866			.4584734	.6185588

Because observations within the same hospital (provnum) are likely to be correlated, we specified the vce(cluster provnum) option. The results show that whether the patient died in the hospital and the type of admission have significant effects on the patient's length of stay.

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Example 2

To illustrate truncated negative binomial regression with more complex data than the previous example, similar data were created from 100 hospitals. Each hospital had its own way of tracking patient data. In particular, hospitals only recorded data from patients with a minimum length of stay, denoted by the variable minstay.

Definitions for minimum length of stay varied among hospitals, typically, from 5 to 18 days. The objective of this example is the same as before: to determine whether the length of stay, recorded in los, was related to the binary variables: died, hmo, type1, type2, and type3.

The binary variables encode the same information as in example 1 above. The minstay variable was used to allow for varying truncation points.

. use https://	/www.stata-pre	ss.com/data	a/r18/med	provider	S	
. tnbreg los d	died hmo type2	-type3, 11((minstay)	vce(clu	ster hospital)	nolog
Truncated negative binomial regression Truncation points: minstay Dispersion: mean Log pseudolikelihood = -7864.0928			Number of obs Wald chi2(4) Prob > chi2 Pseudo R2	s = 2,144 = 15.22 = 0.0043 = 0.0007		
TOP PROMOLING			. adjust	ed for 1	00 clusters in	
		Robust				
los	Coefficient	std. err.	Z	P> z	[95% conf.	interval]
died	.078104	.0303598	2.57	0.010	.0185998	.1376081
hmo	0731132	.0368897	-1.98	0.047	1454158	0008107
type2	.0294132	.0390166	0.75	0.451	047058	.1058845
type3	.0626348	.0540123	1.16	0.246	0432273	.168497
_cons	3.014964	.0290895	103.64	0.000	2.95795	3.071978
/lnalpha	996512	.0828691			-1.158932	8340916
alpha	.3691649	.0305923			.313821	.4342688

In this analysis, two variables have a statistically significant relationship with length of stay. On average, patients who died in the hospital had longer lengths of stay (p = 0.01). Because the coefficient for HMO is negative, that is, $b_{\rm HMO} = -0.073$, on average, patients who were insured by an HMO had shorter lengths of stay (p = 0.047). The type of admission was not statistically significant (p > 0.05).

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Stored results

tnbreg stores the following in e():

Scalars	
e(N)	number of observations
e(k)	number of parameters
e(k_aux)	number of auxiliary parameters
e(k_eq)	number of equations in e(b)
e(k_eq_mode	
e(k_dv)	number of dependent variables
e(df_m)	model degrees of freedom
e(r2_p)	pseudo- R^2
e(11)	log likelihood
e(11_0)	log likelihood, constant-only model
e(ll_c)	log likelihood, comparison model
e(alpha)	value of alpha
e(delta)	value of delta
e(N_clust)	number of clusters
e(chi2)	χ^2
e(chi2_c)	χ^2 for comparison test
e(p)	<i>p</i> -value for model test
e(rank)	rank of e(V)
e(rank0)	rank of e(V) for constant-only model
e(ic)	number of iterations
e(rc)	return code
e(converged) 1 if converged, 0 otherwise
Macros	
e(cmd)	tnbreg
e(cmdline)	command as typed
e(depvar)	name of dependent variable
e(llopt)	contents of 11(), or 0 if not specified
e(wtype)	weight type
e(wexp)	weight expression
e(title)	title in estimation output
e(clustvar)	name of cluster variable
e(offset)	linear offset variable
e(chi2type)	Wald or LR; type of model χ^2 test
e(chi2_ct)	Wald or LR; type of model χ^2 test corresponding to e(chi2_c)
e(dispers)	mean or constant
e(vce)	vcetype specified in vce()
e(vcetype)	title used to label Std. err.
e(opt)	type of optimization
e(which)	max or min; whether optimizer is to perform maximization or minimization
e(ml_method	
e(user)	name of likelihood-evaluator program
e(technique	
e(propertie	
e(predict)	program used to implement predict
e(asbalance	
e(asobserve	d) factor variables fvset as asobserved
Matrices	
e(b)	coefficient vector
e(Cns)	constraints matrix
e(ilog)	iteration log (up to 20 iterations)
e(gradient)	gradient vector
e(V)	variance-covariance matrix of the estimators
e(V_modelba	sed) model-based variance
Functions	
e(sample)	marks estimation sample
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In addition to the above, the following is stored in r():

Matrices r(table)

matrix containing the coefficients with their standard errors, test statistics, *p*-values, and confidence intervals

Note that results stored in r() are updated when the command is replayed and will be replaced when any r-class command is run after the estimation command.

Methods and formulas

Methods and formulas are presented under the following headings:

Mean-dispersion model Constant-dispersion model

Mean-dispersion model

A negative binomial distribution can be regarded as a gamma mixture of Poisson random variables. The number of times an event occurs, y_j , is distributed as $Poisson(\nu_j \mu_j)$. That is, its conditional likelihood is

$$f(y_j \mid \nu_j) = \frac{(\nu_j \mu_j)^{y_j} e^{-\nu_j \mu_j}}{\Gamma(y_j + 1)}$$

where $\mu_j = \exp(\mathbf{x}_j \boldsymbol{\beta} + \text{offset}_j)$ and ν_j is an unobserved parameter with a Gamma $(1/\alpha, \alpha)$ density:

$$g(\nu) = \frac{\nu^{(1-\alpha)/\alpha} e^{-\nu/\alpha}}{\alpha^{1/\alpha} \Gamma(1/\alpha)}$$

This gamma distribution has a mean of 1 and a variance of α , where α is our ancillary parameter.

The unconditional likelihood for the jth observation is therefore

$$f(y_j) = \int_0^\infty f(y_j \mid \nu) g(\nu) \, d\nu = \frac{\Gamma(m+y_j)}{\Gamma(y_j+1)\Gamma(m)} \, p_j^m (1-p_j)^{y_j}$$

where $p_j = 1/(1 + \alpha \mu_j)$ and $m = 1/\alpha$. Solutions for α are handled by searching for $\ln \alpha$ because α must be greater than zero. The conditional probability of observing y_j events given that y_j is greater than the truncation point τ_j is

$$\Pr(Y = y_j \mid y_j > \tau_j, \mathbf{x}_j) = \frac{f(y_j)}{\Pr(Y > \tau_j \mid \mathbf{x}_j)}$$

The log likelihood (with weights w_i and offsets) is given by

$$m = 1/\alpha$$
 $p_j = 1/(1 + \alpha \mu_j)$ $\mu_j = \exp(\mathbf{x}_j \boldsymbol{\beta} + \text{offset}_j)$

$$\ln L = \sum_{j=1}^{n} w_j \left[\ln\{\Gamma(m+y_j)\} - \ln\{\Gamma(y_j+1)\} - \ln\{\Gamma(m)\} + m\ln(p_j) + y_j\ln(1-p_j) - \ln\{\Pr(Y > \tau_j \mid p_j, m)\} \right]$$

Constant-dispersion model

The constant-dispersion model assumes that y_j is conditionally distributed as $Poisson(\mu_j^*)$, where $\mu_j^* \sim Gamma(\mu_j/\delta, \delta)$ for some dispersion parameter δ [by contrast, the mean-dispersion model assumes that $\mu_j^* \sim Gamma(1/\alpha, \alpha \mu_j)$]. The log likelihood is given by

$$m_i = \mu_i / \delta$$
 $p = 1/(1+\delta)$

$$\ln L = \sum_{j=1}^{n} w_j \left[\ln\{\Gamma(m_j + y_j)\} - \ln\{\Gamma(y_j + 1)\} - \ln\{\Gamma(m_j)\} + m_j \ln(p) + y_j \ln(1-p) - \ln\{\Pr(Y > \tau_j \mid p, m_j)\} \right]$$

with everything else defined as shown above in the calculations for the mean-dispersion model.

This command supports the Huber/White/sandwich estimator of the variance and its clustered version using vce(robust) and vce(cluster *clustvar*), respectively. See [P] **_robust**, particularly *Maximum likelihood estimators* and *Methods and formulas*.

tnbreg also supports estimation with survey data. For details on VCEs with survey data, see [SVY] Variance estimation.

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References

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Also see

- [R] tnbreg postestimation Postestimation tools for thbreg
- [R] nbreg Negative binomial regression
- [R] **poisson** Poisson regression
- [R] tpoisson Truncated Poisson regression
- [R] zinb Zero-inflated negative binomial regression
- [R] zip Zero-inflated Poisson regression
- [BAYES] bayes: tnbreg Bayesian truncated negative binomial regression
- [SVY] svy estimation Estimation commands for survey data
- [XT] **xtnbreg** Fixed-effects, random-effects, & population-averaged negative binomial models
- [U] 20 Estimation and postestimation commands

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