

meqrpoisson — Multilevel mixed-effects Poisson regression (QR decomposition)

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Description

`meqrpoisson` is a legacy command for fitting mixed-effects models to binary or binomial responses. `mepoisson` is the modern command, and it offers more functionality; see [ME] [mepoisson](#). The two commands use different but equivalent estimation methods. `mepoisson` performs optimization using variance components in their original metric, whereas `meqrpoisson` uses the QR decomposition of the variance-components matrix.

Quick start

Two-level Poisson regression of `y` on `x` with random intercepts by `lev2` using QR decomposition

```
meqrpoisson y x || lev2:
```

As above, but report incidence-rate ratios

```
meqrpoisson y x || lev2:, irr
```

Add [indicator variables](#) for each level of categorical variable `a` and random coefficients on `x`

```
meqrpoisson y x i.a || lev2: x, irr
```

Three-level random-intercept model of `y` on `x` with `lev2` nested within `lev3`

```
meqrpoisson y x || lev3: || lev2:
```

Menu

Statistics > Multilevel mixed-effects models > Estimation by QR decomposition > Poisson regression

Syntax

```
meqrpoisson depvar fe_equation || re_equation [ || re_equation ... ] [ , options ]
```

where the syntax of *fe_equation* is

```
[ indepvars ] [ if ] [ in ] [ , fe_options ]
```

and the syntax of *re_equation* is one of the following:

for random coefficients and intercepts

```
levelvar: [ varlist ] [ , re_options ]
```

for random effects among the values of a factor variable

```
levelvar: R.varname [ , re_options ]
```

levelvar is a variable identifying the group structure for the random effects at that level or is `_all` representing one group comprising all observations.

<i>fe_options</i>	Description
Model	
<code>noconstant</code>	suppress constant term from the fixed-effects equation
<code>exposure(<i>varname_e</i>)</code>	include $\ln(\text{varname}_e)$ in model with coefficient constrained to 1
<code>offset(<i>varname_o</i>)</code>	include <i>varname_o</i> in model with coefficient constrained to 1

<i>re_options</i>	Description
Model	
<code>covariance(<i>vartype</i>)</code>	variance–covariance structure of the random effects
<code>noconstant</code>	suppress constant term from the random-effects equation
<code>collinear</code>	keep collinear variables

<i>options</i>	Description
Reporting	
<u>level</u> (#)	set confidence level; default is <code>level(95)</code>
<u>irr</u>	report fixed-effects coefficients as incidence-rate ratios
<u>variance</u>	show random-effects parameter estimates as variances and covariances; the default
<u>stddeviations</u>	show random-effects parameter estimates as standard deviations and correlations
<u>noretale</u>	suppress random-effects table
<u>nofetable</u>	suppress fixed-effects table
<u>estmetric</u>	show parameter estimates as stored in <code>e(b)</code>
<u>noheader</u>	suppress output header
<u>nogroup</u>	suppress table summarizing groups
<i>display_options</i>	control columns and column formats, row spacing, line width, display of omitted variables and base and empty cells, and factor-variable labeling
Integration	
<u>intpoints</u> (# [<i># ...</i>])	set the number of integration (quadrature) points; default is <code>intpoints(7)</code>
<u>laplace</u>	use Laplacian approximation; equivalent to <code>intpoints(1)</code>
Maximization	
<i>maximize_options</i>	control the maximization process; seldom used
<u>retolerance</u> (#)	tolerance for random-effects estimates; default is <code>retolerance(1e-8)</code> ; seldom used
<u>reiterate</u> (#)	maximum number of iterations for random-effects estimation; default is <code>reiterate(50)</code> ; seldom used
<u>matsqrt</u>	parameterize variance components using matrix square roots; the default
<u>matlog</u>	parameterize variance components using matrix logarithms
<u>refineopts</u> (<i>maximize_options</i>)	control the maximization process during refinement of starting values
<u>coeflegend</u>	display legend instead of statistics
<hr/>	
<i>vartype</i>	Description
<u>independent</u>	one unique variance parameter per random effect, all covariances 0; the default unless the <code>R.</code> notation is used
<u>exchangeable</u>	equal variances for random effects, and one common pairwise covariance
<u>identity</u>	equal variances for random effects, all covariances 0; the default if the <code>R.</code> notation is used
<u>unstructured</u>	all variances and covariances to be distinctly estimated

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

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

bootstrap, *by*, *jackknife*, *mi estimate*, *rolling*, and *statsby* are allowed; see [U] 11.1.10 Prefix commands.

coeflegend does not appear in the dialog box.

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

Options

Model

noconstant suppresses the constant (intercept) term and may be specified for the fixed-effects equation and for any of or all the random-effects equations.

exposure(*varname_e*) specifies a variable that reflects the amount of exposure over which the *depvar* events were observed for each observation; $\ln(\text{varname}_e)$ is included in the fixed-effects portion of the model with the coefficient constrained to be 1.

offset(*varname_o*) specifies that *varname_o* be included in the fixed-effects portion of the model with the coefficient constrained to be 1.

covariance(*vartype*) specifies the structure of the covariance matrix for the random effects and may be specified for each random-effects equation. *vartype* is one of the following: **independent**, **exchangeable**, **identity**, and **unstructured**.

covariance(**independent**) covariance structure allows for a distinct variance for each random effect within a random-effects equation and assumes that all covariances are 0. The default is **covariance**(**independent**), except when the R. notation is used, in which case the default is **covariance**(**identity**) and only **covariance**(**identity**) and **covariance**(**exchangeable**) are allowed.

covariance(**exchangeable**) structure specifies one common variance for all random effects and one common pairwise covariance.

covariance(**identity**) is short for “multiple of the identity”; that is, all variances are equal and all covariances are 0.

covariance(**unstructured**) allows for all variances and covariances to be distinct. If an equation consists of p random-effects terms, the unstructured covariance matrix will have $p(p+1)/2$ unique parameters.

collinear specifies that **meqrpoisson** not omit collinear variables from the random-effects equation.

Usually, there is no reason to leave collinear variables in place; in fact, doing so usually causes the estimation to fail because of the matrix singularity caused by the collinearity. However, with certain models (for example, a random-effects model with a full set of contrasts), the variables may be collinear, yet the model is fully identified because of restrictions on the random-effects covariance structure. In such cases, using the **collinear** option allows the estimation to take place with the random-effects equation intact.

Reporting

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

irr reports estimated fixed-effects coefficients transformed to incidence-rate ratios, that is, $\exp(\beta)$ rather than β . Standard errors and confidence intervals are similarly transformed. This option affects how results are displayed, not how they are estimated. **irr** may be specified at estimation or upon replay.

variance, the default, displays the random-effects parameter estimates as variances and covariances.

`stddeviations` displays the random-effects parameter estimates as standard deviations and correlations.

`norettable` suppresses the random-effects table.

`nofetable` suppresses the fixed-effects table.

`estmetric` displays all parameter estimates in one table using the metric in which they are stored in `e(b)`. The results are stored in the same metric regardless of the parameterization of the variance components, `matsqrt` or `matlog`, used at estimation time. Random-effects parameter estimates are stored as log-standard deviations and hyperbolic arctangents of correlations, with equation names that organize them by model level. Note that fixed-effects estimates are always stored and displayed in the same metric.

`noheader` suppresses the output header, either at estimation or upon replay.

`nogroup` suppresses the display of group summary information (number of groups, average group size, minimum, and maximum) from the output header.

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

Integration

`intpoints(# [# ...])` sets the number of integration points for adaptive Gaussian quadrature. The more integration points, the more accurate the approximation to the log likelihood. However, computation time increases with the number of quadrature points, and in models with many levels or many random coefficients, this increase can be substantial.

You may specify one number of integration points applying to all levels of random effects in the model, or you may specify distinct numbers of points for each level. `intpoints(7)` is the default; that is, by default seven quadrature points are used for each level.

`laplace` specifies that log likelihoods be calculated using the Laplacian approximation, equivalent to adaptive Gaussian quadrature with one integration point for each level in the model; `laplace` is equivalent to `intpoints(1)`. Computation time increases as a function of the number of quadrature points raised to a power equaling the dimension of the random-effects specification. The computational time saved by using `laplace` can thus be substantial, especially when you have many levels or random coefficients.

The Laplacian approximation has been known to produce biased parameter estimates, but the bias tends to be more prominent in the estimates of the variance components rather than in the estimates of the fixed effects. If your interest lies primarily with the fixed-effects estimates, the Laplace approximation may be a viable faster alternative to adaptive quadrature with multiple integration points.

When the `R.varname` notation is used, the dimension of the random effects increases by the number of distinct values of `varname`. Even when this number is small to moderate, it increases the total random-effects dimension to the point where estimation with more than one quadrature point is prohibitively intensive.

For this reason, when you use the `R.` notation in your random-effects equations, the `laplace` option is assumed. You can override this behavior by using the `intpoints()` option, but doing so is not recommended.

maximize_options: `difficult`, `technique(algorithm_spec)`, `iterate(#)`, `[no]log`, `trace`, `gradient`, `showstep`, `hessian`, `showtolerance`, `tolerance(#)`, `ltolerance(#)`, `nrtolerance(#)`, `nonrtoerance`, and `from(init_specs)`; see [R] [maximize](#). Those that require special mention for `meqrpoisson` are listed below.

For the `technique()` option, the default is `technique(nr)`. The `bhhh` algorithm may not be specified.

`from(init_specs)` is particularly useful when combined with `refineopts(iterate(0))` (see the description [below](#)), which bypasses the initial optimization stage.

`retolerance(#)` specifies the convergence tolerance for the estimated random effects used by adaptive Gaussian quadrature. Although not estimated as model parameters, random-effects estimators are used to adapt the quadrature points. Estimating these random effects is an iterative procedure, with convergence declared when the maximum relative change in the random effects is less than `retolerance()`. The default is `retolerance(1e-8)`. You should seldom have to use this option.

`reiterate(#)` specifies the maximum number of iterations used when estimating the random effects to be used in adapting the Gaussian quadrature points; see the `retolerance()` option. The default is `reiterate(50)`. You should seldom have to use this option.

`matsqrt` (the default), during optimization, parameterizes variance components by using the matrix square roots of the variance–covariance matrices formed by these components at each model level.

`matlog`, during optimization, parameterizes variance components by using the matrix logarithms of the variance–covariance matrices formed by these components at each model level.

The `matsqrt` parameterization ensures that variance–covariance matrices are positive semidefinite, while `matlog` ensures matrices that are positive definite. For most problems, the matrix square root is more stable near the boundary of the parameter space. However, if convergence is problematic, one option may be to try the alternate `matlog` parameterization. When convergence is not an issue, both parameterizations yield equivalent results.

`refineopts(maximize_options)` controls the maximization process during the refinement of starting values. Estimation in `meqrpoisson` takes place in two stages. In the first stage, starting values are refined by holding the quadrature points fixed between iterations. During the second stage, quadrature points are adapted with each evaluation of the log likelihood. Maximization options specified within `refineopts()` control the first stage of optimization; that is, they control the refining of starting values.

maximize_options specified outside `refineopts()` control the second stage.

The one exception to the above rule is the `nolog` option, which when specified outside `refineopts()` applies globally.

`from(init_specs)` is not allowed within `refineopts()` and instead must be specified globally.

Refining starting values helps make the iterations of the second stage (those that lead toward the solution) more numerically stable. In this regard, of particular interest is `refineopts(iterate(#))`, with two iterations being the default. Should the maximization fail because of instability in the Hessian calculations, one possible solution may be to increase the number of iterations here.

The following option is available with `meqrpoisson` but is not shown in the dialog box:

`coeflegend`; see [R] [estimation options](#).

Remarks and examples

Mixed-effects Poisson regression is Poisson regression containing both fixed effects and random effects. In longitudinal data and panel data, random effects are useful for modeling intraclass correlation; that is, observations in the same cluster are correlated because they share common cluster-level random effects.

`meqrpoisson` allows for many levels of random effects. For example, in a three-level model you can specify random effects for schools and then random effects for classes nested within schools. The observations (students, presumably) would comprise level one of the model, the classes would comprise level two, and the schools would comprise level three.

However, for simplicity, for now we consider the two-level model, where for a series of M independent clusters, and conditional on a set of random effects \mathbf{u}_j ,

$$\Pr(y_{ij} = y | \mathbf{u}_j) = \exp(-\mu_{ij}) \mu_{ij}^y / y! \quad (1)$$

for $\mu_{ij} = \exp(\mathbf{x}_{ij}\boldsymbol{\beta} + \mathbf{z}_{ij}\mathbf{u}_j)$, $j = 1, \dots, M$ clusters, and with cluster j consisting of $i = 1, \dots, n_j$ observations. The responses are counts y_{ij} . The $1 \times p$ row vector \mathbf{x}_{ij} are the covariates for the fixed effects, analogous to the covariates you would find in a standard Poisson regression model, with regression coefficients (fixed effects) $\boldsymbol{\beta}$.

The $1 \times q$ vector \mathbf{z}_{ij} are the covariates corresponding to the random effects and can be used to represent both random intercepts and random coefficients. For example, in a random-intercept model, \mathbf{z}_{ij} is simply the scalar 1. The random effects \mathbf{u}_j are M realizations from a multivariate normal distribution with mean $\mathbf{0}$ and $q \times q$ variance matrix $\boldsymbol{\Sigma}$. The random effects are not directly estimated as model parameters but are instead summarized according to the unique elements of $\boldsymbol{\Sigma}$, known as variance components. One special case of (1) places $\mathbf{z}_{ij} = \mathbf{x}_{ij}$ so that all covariate effects are essentially random and distributed as multivariate normal with mean $\boldsymbol{\beta}$ and variance $\boldsymbol{\Sigma}$.

Model (1) is an example of a generalized linear mixed model (GLMM), which generalizes the linear mixed-effects (LME) model to non-Gaussian responses. You can fit LMEs in Stata by using `mixed` and fit GLMMs by using `meglm`. Because of the relationship between LMEs and GLMMs, there is insight to be gained through examination of the linear mixed model. This is especially true for Stata users because the terminology, syntax, options, and output for fitting these types of models are nearly identical. See [ME] `mixed` and the references therein, particularly in the *Introduction*, for more information.

Log-likelihood calculations for fitting any generalized mixed-effects model require integrating out the random effects. One widely used modern method is to directly estimate the integral required to calculate the log likelihood by Gauss–Hermite quadrature or some variation thereof. The estimation method used by `meqrpoisson` is a multicoefficient and multilevel extension of one of these quadrature types, namely, adaptive Gaussian quadrature (AGQ) based on conditional modes, with the multicoefficient extension from [Pinheiro and Bates \(1995\)](#) and the multilevel extension from [Pinheiro and Chao \(2006\)](#); see *Methods and formulas*.

► Example 1: Two-level random-intercept model

[Breslow and Clayton \(1993\)](#) fit a mixed-effects Poisson model to data from a randomized trial of the drug progabide for the treatment of epilepsy.

```
. use http://www.stata-press.com/data/r15/epilepsy
(Epilepsy data; progabide drug treatment)
. describe
Contains data from http://www.stata-press.com/data/r15/epilepsy.dta
  obs:                236                Epilepsy data; progabide drug
                                         treatment
  vars:                8                31 May 2016 14:09
  size:               4,956            (_dta has notes)
```

variable name	storage type	display format	value label	variable label
subject	byte	%9.0g		Subject ID: 1-59
seizures	int	%9.0g		No. of seizures
treat	byte	%9.0g		1: progabide; 0: placebo
visit	float	%9.0g		Dr. visit; coded as (-.3, -.1, .1, .3)
lage	float	%9.0g		log(age), mean-centered
lbas	float	%9.0g		log(0.25*baseline seizures), mean-centered
lbas_trt	float	%9.0g		lbas/treat interaction
v4	byte	%8.0g		Fourth visit indicator

Sorted by: subject

Originally from [Thall and Vail \(1990\)](#), data were collected on 59 subjects (31 on progabide, 28 on placebo). The number of epileptic seizures (`seizures`) was recorded during the two weeks prior to each of four doctor visits (`visit`). The treatment group is identified by the indicator variable `treat`. Data were also collected on the logarithm of age (`lage`) and the logarithm of one-quarter the number of seizures during the eight weeks prior to the study (`lbas`). The variable `lbas_trt` represents the interaction between `lbas` and treatment. `lage`, `lbas`, and `lbas_trt` are mean centered. Because the study originally noted a substantial decrease in seizures prior to the fourth doctor visit, an indicator, `v4`, for the fourth visit was also recorded.

[Breslow and Clayton \(1993\)](#) fit a random-effects Poisson model for the number of observed seizures

$$\log(\mu_{ij}) = \beta_0 + \beta_1 \text{treat}_{ij} + \beta_2 \text{lbas}_{ij} + \beta_3 \text{lbas_trt}_{ij} + \beta_4 \text{lage}_{ij} + \beta_5 \text{v4}_{ij} + u_j$$

for $j = 1, \dots, 59$ subjects and $i = 1, \dots, 4$ visits. The random effects u_j are assumed to be normally distributed with mean 0 and variance σ_u^2 .

```

. meqrpoisson seizures treat lbas lbas_trt lage v4 || subject:
Refining starting values:
Iteration 0:   log likelihood = -680.40577   (not concave)
Iteration 1:   log likelihood = -668.60112
Iteration 2:   log likelihood = -666.37635
Performing gradient-based optimization:
Iteration 0:   log likelihood = -666.37635
Iteration 1:   log likelihood = -665.4589
Iteration 2:   log likelihood = -665.29074
Iteration 3:   log likelihood = -665.29068
Mixed-effects Poisson regression
Group variable: subject
Number of obs      =      236
Number of groups   =       59
Obs per group:
    min =          4
    avg =          4.0
    max =          4
Integration points =    7
Log likelihood     = -665.29068
Wald chi2(5)      =      121.67
Prob > chi2       =      0.0000

```

seizures	Coef.	Std. Err.	z	P> z	[95% Conf. Interval]	
treat	-.9330388	.4008346	-2.33	0.020	-1.71866	-.1474175
lbas	.884433	.1312313	6.74	0.000	.6272244	1.141642
lbas_trt	.3382609	.2033384	1.66	0.096	-.0602752	.7367969
lage	.4842389	.3472775	1.39	0.163	-.1964126	1.16489
v4	-.1610871	.0545758	-2.95	0.003	-.2680537	-.0541206
_cons	2.154574	.2200426	9.79	0.000	1.723299	2.58585

Random-effects Parameters	Estimate	Std. Err.	[95% Conf. Interval]	
subject: Identity				
var(_cons)	.2528265	.058956	.1600785	.3993118

```
LR test vs. Poisson model: chibar2(01) = 304.74      Prob >= chibar2 = 0.0000
```

The number of seizures before the fourth visit does exhibit a significant drop, and the patients on progabide demonstrate a decrease in frequency of seizures compared with the placebo group. The subject-specific random effects also appear significant: $\hat{\sigma}_u^2 = 0.25$ with standard error 0.06.

Because this is a simple random-intercept model, you can obtain equivalent results by using `xtpoisson` with the `re` and `normal` options.

◀

▶ Example 2: Three-level random-intercept model

Rabe-Hesketh and Skrondal (2012, exercise 13.7) describe data from the *Atlas of Cancer Mortality in the European Economic Community* (EEC) (Smans, Mair, and Boyle 1993). The data were analyzed in Langford, Bentham, and McDonald (1998) and record the number of deaths among males due to malignant melanoma during 1971–1980.

```
. use http://www.stata-press.com/data/r15/melanoma
(Skin cancer (melanoma) data)
. describe
Contains data from http://www.stata-press.com/data/r15/melanoma.dta
  obs:          354                Skin cancer (melanoma) data
  vars:          6                  30 May 2016 17:10
  size:         4,956              (_dta has notes)
```

variable name	storage type	display format	value label	variable label
nation	byte	%11.0g	n	Nation ID
region	byte	%9.0g		Region ID: EEC level-I areas
county	int	%9.0g		County ID: EEC level-II/level-III areas
deaths	int	%9.0g		No. deaths during 1971-1980
expected	float	%9.0g		No. expected deaths
uv	float	%9.0g		UV dose, mean-centered

Sorted by:

Nine European nations (variable `nation`) are represented, and data were collected over geographical regions defined by EEC statistical services as level I areas (variable `region`), with deaths being recorded for each of 354 counties, which are level II or level III EEC-defined areas (variable `county`, which identifies the observations). Counties are nested within regions, and regions are nested within nations.

The variable `deaths` records the number of deaths for each county, and `expected` records the expected number of deaths (the exposure) on the basis of crude rates for the combined countries. Finally, the variable `uv` is a measure of exposure to ultraviolet (UV) radiation.

In modeling the number of deaths, one possibility is to include dummy variables for the nine nations as fixed effects. Another is to treat these as random effects and fit the three-level random-intercept Poisson model,

$$\log(\mu_{ijk}) = \log(\text{expected}_{ijk}) + \beta_0 + \beta_1 \text{uv}_{ijk} + \beta_2 \text{uv}_{ijk}^2 + u_k + v_{jk}$$

for nation k , region j , and county i . The model includes an exposure term for expected deaths.

```
. meqrpoisson deaths c.uv##c.uv, exposure(expected) || nation: || region:
Refining starting values:
Iteration 0: log likelihood = -1169.4088 (not concave)
Iteration 1: log likelihood = -1156.8957 (not concave)
Iteration 2: log likelihood = -1101.8457
Performing gradient-based optimization:
Iteration 0: log likelihood = -1101.8457
Iteration 1: log likelihood = -1090.6214
Iteration 2: log likelihood = -1089.4198
Iteration 3: log likelihood = -1089.411
Iteration 4: log likelihood = -1089.411
Mixed-effects Poisson regression          Number of obs    =      354
```

Group Variable	No. of Groups	Observations per Group			Integration Points
		Minimum	Average	Maximum	
nation	9	3	39.3	95	7
region	78	1	4.5	13	7

```
Log likelihood = -1089.411          Wald chi2(2)    =      25.69
                                   Prob > chi2       =      0.0000
```

deaths	Coef.	Std. Err.	z	P> z	[95% Conf. Interval]	
uv	.0056975	.0137931	0.41	0.680	-.0213364	.0327314
c.uv#c.uv	-.0058374	.001388	-4.21	0.000	-.0085579	-.0031169
_cons ln(expected)	.1289976 1	.1581122 (exposure)	0.82	0.415	-.1808967	.4388918

Random-effects Parameters		Estimate	Std. Err.	[95% Conf. Interval]	
nation: Identity	var(_cons)	.1840722	.094531	.0672745	.5036466
region: Identity	var(_cons)	.0382743	.0087869	.0244057	.0600237

```
LR test vs. Poisson model: chi2(2) = 1267.13          Prob > chi2 = 0.0000
```

Note: LR test is conservative and provided only for reference.

By including an exposure variable that is an expected rate, we are in effect specifying a linear model for the log of the standardized mortality ratio, the ratio of observed deaths to expected deaths that is based on a reference population. Here the reference population is all nine nations.

Looking at the estimated variance components, we can see that there is more unexplained variability between nations than between regions within each nation. This may be due to, for example, country-specific informational campaigns on the risks of sun exposure.

Stored results

meqrpoisson stores the following in `e()`:

Scalars

<code>e(N)</code>	number of observations
<code>e(k)</code>	number of parameters
<code>e(k_f)</code>	number of fixed-effects parameters
<code>e(k_r)</code>	number of random-effects parameters
<code>e(k_rs)</code>	number of variances
<code>e(k_rc)</code>	number of covariances
<code>e(df_m)</code>	model degrees of freedom
<code>e(ll)</code>	log likelihood
<code>e(chi2)</code>	χ^2
<code>e(p)</code>	p -value for model test
<code>e(ll_c)</code>	log likelihood, comparison model
<code>e(chi2_c)</code>	χ^2 , comparison test
<code>e(df_c)</code>	degrees of freedom, comparison test
<code>e(p_c)</code>	p -value for comparison test
<code>e(rank)</code>	rank of <code>e(V)</code>
<code>e(ic)</code>	number of iterations
<code>e(reparam_rc)</code>	return code, final reparameterization
<code>e(rc)</code>	return code
<code>e(converged)</code>	1 if converged, 0 otherwise

Macros

<code>e(cmd)</code>	meqrpoisson
<code>e(cmdline)</code>	command as typed
<code>e(depvar)</code>	name of dependent variable
<code>e(ivars)</code>	grouping variables
<code>e(exposurevar)</code>	exposure variable
<code>e(model)</code>	Poisson
<code>e(title)</code>	title in estimation output
<code>e(offset)</code>	offset
<code>e(redim)</code>	random-effects dimensions
<code>e(vartypes)</code>	variance-structure types
<code>e(revars)</code>	random-effects covariates
<code>e(n_quad)</code>	number of integration points
<code>e(laplace)</code>	laplace, if Laplace approximation
<code>e(chi2type)</code>	Wald; type of model χ^2
<code>e(method)</code>	ML
<code>e(opt)</code>	type of optimization
<code>e(ml_method)</code>	type of ml method
<code>e(technique)</code>	maximization technique
<code>e(datasignature)</code>	the checksum
<code>e(datasignaturevars)</code>	variables used in calculation of checksum
<code>e(properties)</code>	b V
<code>e(estat_cmd)</code>	program used to implement estat
<code>e(predict)</code>	program used to implement predict
<code>e(marginsnotok)</code>	predictions disallowed by margins
<code>e(marginsdefault)</code>	default predict() specification for margins
<code>e(asbalanced)</code>	factor variables fvset as asbalanced
<code>e(asobserved)</code>	factor variables fvset as asobserved

Matrices

<code>e(b)</code>	coefficient vector
<code>e(N_g)</code>	group counts
<code>e(g_min)</code>	group-size minimums
<code>e(g_avg)</code>	group-size averages
<code>e(g_max)</code>	group-size maximums
<code>e(V)</code>	variance-covariance matrix of the estimators

Functions

<code>e(sample)</code>	marks estimation sample
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Methods and formulas

In a two-level Poisson model, for cluster j , $j = 1, \dots, M$, the conditional distribution of $\mathbf{y}_j = (y_{j1}, \dots, y_{jn_j})'$, given a set of cluster-level random effects \mathbf{u}_j , is

$$\begin{aligned} f(\mathbf{y}_j | \mathbf{u}_j) &= \prod_{i=1}^{n_j} [\{\exp(\mathbf{x}_{ij}\boldsymbol{\beta} + \mathbf{z}_{ij}\mathbf{u}_j)\}^{y_{ij}} \exp\{-\exp(\mathbf{x}_{ij}\boldsymbol{\beta} + \mathbf{z}_{ij}\mathbf{u}_j)\} / y_{ij}!] \\ &= \exp \left[\sum_{i=1}^{n_j} \{y_{ij}(\mathbf{x}_{ij}\boldsymbol{\beta} + \mathbf{z}_{ij}\mathbf{u}_j) - \exp(\mathbf{x}_{ij}\boldsymbol{\beta} + \mathbf{z}_{ij}\mathbf{u}_j) - \log(y_{ij}!)\} \right] \end{aligned}$$

Defining $c(\mathbf{y}_j) = \sum_{i=1}^{n_j} \log(y_{ij}!)$, where $c(\mathbf{y}_j)$ does not depend on the model parameters, we can express the above compactly in matrix notation,

$$f(\mathbf{y}_j | \mathbf{u}_j) = \exp \{ \mathbf{y}'_j (\mathbf{X}_j \boldsymbol{\beta} + \mathbf{Z}_j \mathbf{u}_j) - \mathbf{1}' \exp(\mathbf{X}_j \boldsymbol{\beta} + \mathbf{Z}_j \mathbf{u}_j) - c(\mathbf{y}_j) \}$$

where \mathbf{X}_j is formed by stacking the row vectors \mathbf{x}_{ij} and \mathbf{Z}_j is formed by stacking the row vectors \mathbf{z}_{ij} . We extend the definition of $\exp(\cdot)$ to be a vector function where necessary.

Because the prior distribution of \mathbf{u}_j is multivariate normal with mean $\mathbf{0}$ and $q \times q$ variance matrix $\boldsymbol{\Sigma}$, the likelihood contribution for the j th cluster is obtained by integrating \mathbf{u}_j out of the joint density $f(\mathbf{y}_j, \mathbf{u}_j)$,

$$\begin{aligned} \mathcal{L}_j(\boldsymbol{\beta}, \boldsymbol{\Sigma}) &= (2\pi)^{-q/2} |\boldsymbol{\Sigma}|^{-1/2} \int f(\mathbf{y}_j | \mathbf{u}_j) \exp(-\mathbf{u}'_j \boldsymbol{\Sigma}^{-1} \mathbf{u}_j / 2) d\mathbf{u}_j \\ &= \exp\{-c(\mathbf{y}_j)\} (2\pi)^{-q/2} |\boldsymbol{\Sigma}|^{-1/2} \int \exp\{h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j)\} d\mathbf{u}_j \end{aligned} \quad (2)$$

where

$$h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j) = \mathbf{y}'_j (\mathbf{X}_j \boldsymbol{\beta} + \mathbf{Z}_j \mathbf{u}_j) - \mathbf{1}' \exp(\mathbf{X}_j \boldsymbol{\beta} + \mathbf{Z}_j \mathbf{u}_j) - \mathbf{u}'_j \boldsymbol{\Sigma}^{-1} \mathbf{u}_j / 2$$

and for convenience, in the arguments of $h(\cdot)$ we suppress the dependence on the observable data $(\mathbf{y}_j, \mathbf{X}_j, \mathbf{Z}_j)$.

The integration in (2) has no closed form and thus must be approximated. The Laplacian approximation (Tierney and Kadane 1986; Pinheiro and Bates 1995) is based on a second-order Taylor expansion of $h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j)$ about the value of \mathbf{u}_j that maximizes it. Taking first and second derivatives, we obtain

$$\begin{aligned} h'(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j) &= \frac{\partial h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j)}{\partial \mathbf{u}_j} = \mathbf{Z}'_j \{ \mathbf{y}_j - \mathbf{m}(\boldsymbol{\beta}, \mathbf{u}_j) \} - \boldsymbol{\Sigma}^{-1} \mathbf{u}_j \\ h''(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j) &= \frac{\partial^2 h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j)}{\partial \mathbf{u}_j \partial \mathbf{u}'_j} = - \{ \mathbf{Z}'_j \mathbf{V}(\boldsymbol{\beta}, \mathbf{u}_j) \mathbf{Z}_j + \boldsymbol{\Sigma}^{-1} \} \end{aligned}$$

where $\mathbf{m}(\boldsymbol{\beta}, \mathbf{u}_j)$ is the vector function with the i th element equal to the conditional mean of y_{ij} given \mathbf{u}_j , that is, $\exp(\mathbf{x}_{ij}\boldsymbol{\beta} + \mathbf{z}_{ij}\mathbf{u}_j)$. $\mathbf{V}(\boldsymbol{\beta}, \mathbf{u}_j)$ is the diagonal matrix whose diagonal entries v_{ij} are the conditional variances of y_{ij} given \mathbf{u}_j , namely,

$$v_{ij} = \exp(\mathbf{x}_{ij}\boldsymbol{\beta} + \mathbf{z}_{ij}\mathbf{u}_j)$$

because equality of mean and variance is a characteristic of the Poisson distribution.

The maximizer of $h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j)$ is $\hat{\mathbf{u}}_j$ such that $h'(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j) = \mathbf{0}$. The integrand in (2) is proportional to the posterior density $f(\mathbf{u}_j | \mathbf{y}_j)$, so $\hat{\mathbf{u}}_j$ also represents the posterior mode, a plausible estimator of \mathbf{u}_j in its own right.

Given the above derivatives, the second-order Taylor approximation then takes the form

$$h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j) \approx h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j) + \frac{1}{2} (\mathbf{u}_j - \hat{\mathbf{u}}_j)' h''(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j) (\mathbf{u}_j - \hat{\mathbf{u}}_j) \quad (3)$$

The first-derivative term vanishes because $h'(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j) = \mathbf{0}$. Therefore,

$$\begin{aligned} \int \exp\{h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j)\} d\mathbf{u}_j &\approx \exp\{h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j)\} \\ &\quad \times \int \exp\left[-\frac{1}{2} (\mathbf{u}_j - \hat{\mathbf{u}}_j)' \{-h''(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j)\} (\mathbf{u}_j - \hat{\mathbf{u}}_j)\right] d\mathbf{u}_j \quad (4) \\ &= \exp\{h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j)\} (2\pi)^{q/2} |-h''(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j)|^{-1/2} \end{aligned}$$

because the latter integrand can be recognized as the “kernel” of a multivariate normal density.

Combining the above with (2) (and taking logs) gives the Laplacian log-likelihood contribution of the j th cluster,

$$\mathcal{L}_j^{\text{Lap}}(\boldsymbol{\beta}, \boldsymbol{\Sigma}) = -\frac{1}{2} \log |\boldsymbol{\Sigma}| - \log |\mathbf{R}_j| + h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j) - c(\mathbf{y}_j)$$

where \mathbf{R}_j is an upper-triangular matrix such that $-h''(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j) = \mathbf{R}_j \mathbf{R}_j'$. [Pinheiro and Chao \(2006\)](#) show that $\hat{\mathbf{u}}_j$ and \mathbf{R}_j can be efficiently computed as the iterative solution to a least-squares problem by using matrix decomposition methods similar to those used in fitting LME models ([Bates and Pinheiro 1998](#); [Pinheiro and Bates 2000](#); [ME] [mixed](#)).

The fidelity of the Laplacian approximation is determined wholly by the accuracy of the approximation in (3). An alternative that does not depend so heavily on this approximation is integration via AGQ ([Naylor and Smith 1982](#); [Liu and Pierce 1994](#)).

The application of AGQ to this particular problem is from [Pinheiro and Bates \(1995\)](#). When we reexamine the integral in question, a transformation of integration variables yields

$$\begin{aligned} \int \exp\{h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j)\} d\mathbf{u}_j &= |\mathbf{R}_j|^{-1} \int \exp\{h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j + \mathbf{R}_j^{-1} \mathbf{t})\} dt \\ &= (2\pi)^{q/2} |\mathbf{R}_j|^{-1} \int \exp\{h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j + \mathbf{R}_j^{-1} \mathbf{t}) + \mathbf{t}' \mathbf{t} / 2\} \phi(\mathbf{t}) dt \quad (5) \end{aligned}$$

where $\phi(\cdot)$ is the standard multivariate normal density. Because the integrand is now expressed as some function multiplied by a normal density, it can be estimated by applying the rules of standard Gauss–Hermite quadrature. For a predetermined number of quadrature points N_Q , define $a_k = \sqrt{2} a_k^*$ and $w_k = w_k^* / \sqrt{\pi}$, for $k = 1, \dots, N_Q$, where (a_k^*, w_k^*) are a set of abscissas and weights for Gauss–Hermite quadrature approximations of $\int \exp(-x^2) f(x) dx$, as obtained from [Abramowitz and Stegun \(1964, 924\)](#).

Define $\mathbf{a}_k = (a_{k_1}, a_{k_2}, \dots, a_{k_q})'$; that is, \mathbf{a}_k is a vector that spans the N_Q abscissas over the dimension q of the random effects. Applying quadrature rules to (5) yields the AGQ approximation,

$$\begin{aligned} & \int \exp \{h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \mathbf{u}_j)\} d\mathbf{u}_j \\ & \approx (2\pi)^{q/2} |\mathbf{R}_j|^{-1} \sum_{k_1=1}^{N_Q} \cdots \sum_{k_q=1}^{N_Q} \left[\exp \{h(\boldsymbol{\beta}, \boldsymbol{\Sigma}, \hat{\mathbf{u}}_j + \mathbf{R}_j^{-1} \mathbf{a}_k) + \mathbf{a}'_k \mathbf{a}_k / 2\} \prod_{p=1}^q w_{k_p} \right] \\ & \equiv (2\pi)^{q/2} \hat{G}_j(\boldsymbol{\beta}, \boldsymbol{\Sigma}) \end{aligned}$$

resulting in the AGQ log-likelihood contribution of the j th cluster,

$$\mathcal{L}_j^{\text{AGQ}}(\boldsymbol{\beta}, \boldsymbol{\Sigma}) = -\frac{1}{2} \log |\boldsymbol{\Sigma}| + \log \left\{ \hat{G}_j(\boldsymbol{\beta}, \boldsymbol{\Sigma}) \right\} - c(\mathbf{y}_j)$$

The “adaptive” part of adaptive Gaussian quadrature lies in the translation and rescaling of the integration variables in (5) by using $\hat{\mathbf{u}}_j$ and \mathbf{R}_j^{-1} , respectively. This transformation of quadrature abscissas (centered at 0 in standard form) is chosen to better capture the features of the integrand, through which (4) can be seen to resemble a multivariate normal distribution with mean $\hat{\mathbf{u}}_j$ and variance $\mathbf{R}_j^{-1} \mathbf{R}_j^{-T}$. AGQ is therefore not as dependent as the Laplace method upon the approximation in (3). In AGQ, (3) serves merely to redirect the quadrature abscissas, with the AGQ approximation improving as the number of quadrature points, N_Q , increases. In fact, [Pinheiro and Bates \(1995\)](#) point out that AGQ with only one quadrature point ($a = 0$ and $w = 1$) reduces to the Laplacian approximation.

The log likelihood for the entire dataset is then simply the sum of the contributions of the M individual clusters, namely, $\mathcal{L}(\boldsymbol{\beta}, \boldsymbol{\Sigma}) = \sum_{j=1}^M \mathcal{L}_j^{\text{Lap}}(\boldsymbol{\beta}, \boldsymbol{\Sigma})$ for Laplace and $\mathcal{L}(\boldsymbol{\beta}, \boldsymbol{\Sigma}) = \sum_{j=1}^M \mathcal{L}_j^{\text{AGQ}}(\boldsymbol{\beta}, \boldsymbol{\Sigma})$ for AGQ.

Maximization of $\mathcal{L}(\boldsymbol{\beta}, \boldsymbol{\Sigma})$ is performed with respect to $(\boldsymbol{\beta}, \boldsymbol{\theta})$, where $\boldsymbol{\theta}$ is a vector comprising the unique elements of the matrix square root of $\boldsymbol{\Sigma}$. This is done to ensure that $\boldsymbol{\Sigma}$ is always positive semidefinite. If the `matlog` option is specified, then $\boldsymbol{\theta}$ instead consists of the unique elements of the matrix logarithm of $\boldsymbol{\Sigma}$. For well-conditioned problems, both methods produce equivalent results, yet our experience deems the former as more numerically stable near the boundary of the parameter space.

Once maximization is achieved, parameter estimates are mapped from $(\hat{\boldsymbol{\beta}}, \hat{\boldsymbol{\theta}})$ to $(\hat{\boldsymbol{\beta}}, \hat{\boldsymbol{\gamma}})$, where $\hat{\boldsymbol{\gamma}}$ is a vector containing the unique (estimated) elements of $\boldsymbol{\Sigma}$, expressed as logarithms of standard deviations for the diagonal elements and hyperbolic arctangents of the correlations for off-diagonal elements. This last step is necessary to (a) obtain a parameterization under which parameter estimates can be displayed and interpreted individually, rather than as elements of a matrix square root (or logarithm), and (b) parameterize these elements such that their ranges each encompass the entire real line.

Parameter estimates are stored in `e(b)` as $(\hat{\boldsymbol{\beta}}, \hat{\boldsymbol{\gamma}})$, with the corresponding variance–covariance matrix stored in `e(V)`. Parameter estimates can be displayed in this metric by specifying the `estmetric` option. However, in `meqrpoisson` output, variance components are displayed as variances and covariances.

The approach outlined above can be extended from two-level models to models with three or more levels; see [Pinheiro and Chao \(2006\)](#) for details.

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Also see

[ME] **meqrpoisson postestimation** — Postestimation tools for meqrpoisson

[ME] **menbreg** — Multilevel mixed-effects negative binomial regression

[ME] **mepoisson** — Multilevel mixed-effects Poisson regression

[ME] **me** — Introduction to multilevel mixed-effects models

[MI] **estimation** — Estimation commands for use with mi estimate

[SEM] **intro 5** — Tour of models (*Multilevel mixed-effects models*)

[XT] **xtpoisson** — Fixed-effects, random-effects, and population-averaged Poisson models

[U] **20 Estimation and postestimation commands**