Package ‘Countr’

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Type Package
Title Flexible Univariate Count Models Based on Renewal Processes
Version 3.5.4
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Description Flexible univariate count models based on renewal processes. The models may include covariates and can be specified with familiar formula syntax as in glm() and package ‘flexsurv’. The methodology is described in a forthcoming paper in the Journal of Statistical Software <10.18637/jss.v090.i13>(included as vignette ‘Countr_guide’ in the package).

License GPL (>= 2)

URL https://github.com/GeoBosh/Countr
     https://geobosh.github.io/Countr/

BugReports https://github.com/GeoBosh/Countr/issues

Depends R (>= 3.3.0)
Imports Matrix, Rcpp (>= 0.11.3), flexsurv, Formula, VGAM, optimx, numDeriv, boot, MASS, car, utils, Rdpack (>= 0.7-0), lattice, RColorBrewer, dplyr, standardize, pscl, lmtest, xtable

LinkingTo Rcpp, RcppArmadillo

Suggests testthat, knitr

RdMacros Rdpack

VignetteBuilder knitr

LazyData true

RoxygenNote 6.1.1

NeedsCompilation yes

Collate 'Countr-package.R' 'RcppExports.R' 'anc.R' 'coefficient.R'
     'convCount_loglik.R' 'convCount_moments.R' 'convCount_probs.R'
     'dWeibull.R' 'dWeibullgamma.R' 'data.R' 'renewal_IV.R'
     'renewal_tools.R' 'renewal_cstr.R' 'tools.R'
     'renewal_methods.R'
Description

Flexible univariate count models based on renewal processes. The models may include covariates and can be specified with familiar formula syntax as in `glm()` and ‘flexsurv’.
addBootSampleObject

Details

The methodology is described in the forthcoming paper (Kharrat, Boshnakov, McHale, and Baker 2019) in the Journal of Statistical Software (included in the package as vignette vignette("Countr_guide_paper",package = "Countr")).

The main function is renewalCount, see its documentation for examples.

Goodness of fit chi-square (likelihood ratio and Pearson) tests for glm and count renewal models are implemented in chiSq_gof and chiSq_pearson.

References


addBootSampleObject Create a bootstrap sample for coefficient estimates

Description

Create a bootstrap sample from coefficient estimates.

Usage

addBootSampleObject(object, ...)

Arguments

object

an object to add boot object to

... extra parameters to be passed to the boot::boot() function other than data and statistic.
Details

The information in object is used to prepare the arguments and then boot is called to generate the bootstrap sample. The bootstrap sample is stored in object as component "boot". Arguments in "..." can be used customise the boot() call.

Value

object with additional component "boot"

See Also

renewal_methods

Examples

## see renewal_methods

---

chiSq_gof

Formal Chi-square goodness-of-fit test

Description

Carry out the formal chi-square goodness-of-fit test described by Cameron (2013).

Usage

chiSq_gof(object, breaks, ...)

## S3 method for class 'renewal'
chiSq_gof(object, breaks, ...)

## S3 method for class 'negbin'
chiSq_gof(object, breaks, ...)

## S3 method for class 'glm'
chiSq_gof(object, breaks, ...)

Arguments

object an object from class renewal
breaks integer values at which the breaks should happen. The function will compute the observed frequencies in the intervals [breaks[i],breaks[i + 1])
... currently not used
Details

The test is a conditional moment test described in details in Cameron (2013, Section 5.3.4). We compute the asymptotically equivalent outer product of the gradient version which is justified for renewal models (fully parametric + parameters based on MLE).

Value
data.frame

References


See Also

chiSq_pearson

---

## chiSq_pearson

### Pearson Chi-Square test

### Description

Carry out Pearson Chi-Square test and compute the Pearson statistic.

### Usage

```r
chiSq_pearson(object, ...)
```

```r
## S3 method for class 'renewal'
chiSq_pearson(object, ...)
```

```r
## S3 method for class 'glm'
chiSq_pearson(object, ...)
```

### Arguments

- **object**: an object from class renewal
- **...**: currently not used

### Details

The computation is inspired from Cameron(2013) Chapter 5.3.4. Observed and fitted frequencies are computed and the contribution of every observed cell to the Pearson’s chi-square test statistic is reported. The idea is to check if the fitted model has a tendency to over or under predict some ranges of data.
Value
data.frame with 5 columns given the count values (Counts), observed frequencies (Actual), model’s prediction (Predicted), the difference (Diff) and the contribution to the Pearson’s statistic (Pearson).

References

See Also
chiSq_gof

compareToGLM(poisson_model, breaks, nbinom_model, ...)

Arguments
  poisson_model  fitted Poisson glm model
  breaks         integer values at which the breaks should happen. The function will compute the observed frequencies in the intervals [breaks[i],breaks[i + 1])
  nbinom_model   fitted negative binomial (fitted using MASS::glm.nb()). This argument is optional.
  ...            renewal models to be considered.

Details
This function computes a data.frame similar to Table 5.6 in Cameron(2013), using the observed frequencies and predictions from different models. Supported models accepted are Poisson and negative binomial (fitted using MASS::glm.nb()) from the glm family and any model from the renewal family (passed in ...).

Value
data.frame with columns Counts, Actual (observed probability) and then 2 columns per model passed (predicted probability and pearson statistic) for the associated count value.
References


---

Create a formula for renewalCount

**Description**

Create a formula for renewalCount

**Usage**

CountrFormula(response, ...)

**Arguments**

- `response` the formula for the "main" parameter. It also specifies the response variable.
- `...` additional arguments for the ancilliary parameters.

**Value**

a Formula object suitable for argument formula of renewalCount().

---

Summary of a count variable

**Description**

Summary of a count variable.

**Usage**

count_table(count, breaks, formatChar = FALSE)

**Arguments**

- `count` integer, observed count value for every individual in the sample.
- `breaks` integer, values at which the breaks should happen. The function will compute the observed frequency in [breaks[i],breaks[i + 1]).
- `formatChar` logical, should the values be converted to character and formatted?
Details

The function does a similar job to \texttt{table()} with more flexibility introduced by the argument \texttt{breaks}. The user can decide how to break the count values and decide to merge some cells if needed.

Value

\texttt{matrix} with 2 rows and \texttt{length(breaks)} columns. The column names are the cells names. The rows are the observed frequencies and relative frequencies (probabilities).

\begin{verbatim}
dCount_conv_bi
\end{verbatim}

\textit{Compute count probabilities using convolution}

Description

Compute count probabilities using one of several convolution methods. \texttt{dCount_conv_bi} does the computations for the distributions with built-in support in this package. \texttt{dCount_conv_user} does the same using a user defined survival function.

Usage

\begin{verbatim}
dCount_conv_bi(x, distPars, dist = c("weibull", "gamma", "gengamma", "burr"), method = c("dePril", "direct", "naive"), nsteps = 100, time = 1, extrap = TRUE, log = FALSE)
dCount_conv_user(x, distPars, extrapolPars, survR, method = c("dePril", "direct", "naive"), nsteps = 100, time = 1, extrap = TRUE, log = FALSE)
\end{verbatim}

Arguments

\begin{verbatim}
x distPars dist method nsteps time extrap log extrapolPars survR
\end{verbatim}

integer (vector), the desired count values.

\texttt{Rcpp::List} with distribution specific slots, see details.

character name of the built-in distribution, see details.

character string, the method to use, see Details.

unsiged integer, number of steps used to compute the integral.

double, time at which to compute the probabilities. Set to 1 by default.

logical, if \texttt{TRUE}, Richardson extrapolation will be applied to improve accuracy.

logical, if \texttt{TRUE} the log-probability will be returned.

vector of length 2, the extrapolation values.

function, user supplied survival function; should have signature function(\texttt{t, distPars}), where \texttt{t} is a positive real number (the time where the survival function is evaluated) and \texttt{distPars} is a list of distribution parameters. It should return a double value.
**Details**

dCount_conv_bi computes count probabilities using one of several convolution methods for the distributions with built-in support in this package.

The following convolution methods are implemented: "dePril", "direct", and "naive".

The built-in distributions currently are Weibull, gamma, generalised gamma, and Burr.

**Value**

vector of probabilities $P(x(i), i = 1, ..., n)$ where $n$ is the length of $x$.

**Examples**

```r
x <- 0:10
lambda <- 2.56
p0 <- dpois(x, lambda)
ll <- sum(dpois(x, lambda, TRUE))

err <- 1e-6
## all-probs convolution approach
distPars <- list(scale = lambda, shape = 1)
pmat_bi <- dCount_conv_bi(x, distPars, "weibull", "direct",
nsteps = 200)

## user pwei
pwei_user <- function(tt, distP) {
  alpha <- exp(-log(distP["scale"])/distP["shape"])
  pweibull(q = tt, scale = alpha, shape = distP["shape"],
  lower.tail = FALSE)
}

pmat_user <- dCount_conv_user(x, distPars, c(1, 2), pwei_user, "direct",
nsteps = 200)

max((pmat_bi - p0)^2 / p0)
max((pmat_user - p0)^2 / p0)

## naive convolution approach
pmat_bi <- dCount_conv_bi(x, distPars, "weibull", "naive",
nsteps = 200)
pmat_user <- dCount_conv_user(x, distPars, c(1, 2), pwei_user, "naive",
nsteps = 200)

max((pmat_bi - p0)^2 / p0)
max((pmat_user - p0)^2 / p0)

## dePril conv approach
pmat_bi <- dCount_conv_bi(x, distPars, "weibull", "dePril",
nsteps = 200)
pmat_user <- dCount_conv_user(x, distPars, c(1, 2), pwei_user, "dePril",
nsteps = 200)

max((pmat_bi - p0)^2 / p0)
max((pmat_user - p0)^2 / p0)
```
dCount_conv_loglik_bi  

Log-likelihood of a count probability computed by convolution (bi)

Description

Compute the log-likelihood of a count model using convolution methods to compute the probabilities. dCount_conv_loglik_bi is for the built-in distributions. dCount_conv_loglik_user is for user-defined survival functions.

Usage

```r
# Built-in distribution
dCount_conv_loglik_bi(x, distPars, dist = c("weibull", "gamma", "gengamma", "burr"), method = c("dePril", "direct", "naive"), nsteps = 100, time = 1, extrap = TRUE, na.rm = TRUE, weights = NULL)

# User-defined survival functions
dCount_conv_loglik_user(x, distPars, extrapolPars, survR, method = c("dePril", "direct", "naive"), nsteps = 100, time = 1, extrap = TRUE, na.rm = TRUE, weights = NULL)
```

Arguments

- `x` integer (vector), the desired count values.
- `distPars` list of the same length as `x` with each slot being itself a named list containing the distribution parameters corresponding to `x[i]`.
- `dist` character name of the built-in distribution, see details.
- `method` character, convolution method to be used; choices are "dePril" (section 3.2), "direct" (section 2) or "naive" (section 3.1).
- `nsteps` unsigned integer number of steps used to compute the integral.
- `time` double time at which to compute the probabilities. Set to 1 by default.
- `extrap` logical if TRUE, Richardson extrapolation will be applied to improve accuracy.
- `na.rm` logical, if TRUE, NAs (produced by taking the log of very small probabilities) will be replaced by the smallest allowed probability; default is TRUE.
- `weights` numeric, vector of weights to apply. If NULL, a vector of ones.
- `extrapolPars` list of same length as `x` where each slot is a vector of length 2 (the extrapolation values to be used) corresponding to `x[i]`.
- `survR` a user-defined survival function; should have the signature `function(t, distPars)` where `t` is a real number (>0) where the survival function is evaluated and `distPars` is a list of distribution parameters. It should return a double value.

Value

numeric, the log-likelihood of the count process
Examples

```r
x <- 0:10
lambda <- 2.56
distPars <- list(scale = lambda, shape = 1)
distParsList <- lapply(seq(along = x), function(ind) distPars)
extrapolParsList <- lapply(seq(along = x), function(ind) c(2, 1))
## user pwei
pwei_user <- function(tt, distP) {
  alpha <- exp(-log(distP["scale"]) / distP["shape"])
pweibull(q = tt, scale = alpha, shape = distP["shape"],
          lower.tail = FALSE)
}
## log-likelihood allProbs Poisson
dCount_conv_loglik_bi(x, distParsList,
                      "weibull", "direct", nsteps = 400)

dCount_conv_loglik_user(x, distParsList, extrapolParsList,
                        pwei_user, "direct", nsteps = 400)
## log-likelihood naive Poisson

dCount_conv_loglik_bi(x, distParsList,
                      "weibull", "naive", nsteps = 400)

dCount_conv_loglik_user(x, distParsList, extrapolParsList,
                        pwei_user, "naive", nsteps = 400)
## log-likelihood dePril Poisson

dCount_conv_loglik_bi(x, distParsList,
                      "weibull", "dePril", nsteps = 400)

dCount_conv_loglik_user(x, distParsList, extrapolParsList,
                        pwei_user, "dePril", nsteps = 400)
## see dCount_conv_loglik_bi()
```

---

**dmodifiedCount_bi**

*Compute count probabilities based on modified renewal process (bi)*

**Description**

Compute count probabilities based on modified renewal process using dePril algorithm. `dmodifiedCount_bi` does it for the builtin distributions. `dmodifiedCount_user` does the same for a user specified distribution.

**Usage**

```r
dmodifiedCount_bi(x, distPars, dist, distPars0, dist0, nsteps = 100L,
                   time = 1, extrap = TRUE, cdfout = FALSE, logFlag = FALSE)
```
dmodifiedCount_user(x, distPars, survR, distPars0, survR0, extrapolPars,
   nsteps = 100L, time = 1, extrap = TRUE, cdfout = FALSE,
   logFlag = FALSE)

Arguments

x integer (vector), the desired count values.
distPars0, distPars Rcpp::List with distribution specific slots for the first arrival and the rest of the process respectively.
dist0, dist character, name of the first and following survival distributions.
nsteps unsigned integer number of steps used to compute the integral.
time double time at which to compute the probabilities. Set to 1 by default.
extrap logical if TRUE, Richardson extrapolation will be applied to improve accuracy.
cdfout TODO
logFlag logical if TRUE the log-probability will be returned.
survR0, survR user supplied survival function; should have signature function(t,distPars), where t is a positive real number (the time at which the survival function is evaluated) and distPars is a list of distribution parameters. It should return a double value (first arrival and following arrivals respectively).
extrapolPars list of same length as x, where each slot is a vector of length 2 (the extrapolation values to be used) corresponding to x[i].

Details

For the modified renewal process the first arrival is allowed to have a different distribution from the time between subsequent arrivals. The renewal assumption is kept.

Value

vector of probabilities P(x(i)) for i = 1, ..., n where n is the length of x.

---

dWeibullCount Probability calculations for Weibull count models

Description

Probability computations for the univariate Weibull count process. Several methods are provided.
dWeibullCount computes probabilities.
dWeibullCount_loglik computes the log-likelihood.
evWeibullCount computes the expected value and variance.
Usage

dWeibullCount(x, shape, scale, method = c("series_acc", "series_mat", "conv_direct", "conv_naive", "conv_dePril"), time = 1, log = FALSE, conv_steps = 100, conv_extrap = TRUE, series_terms = 50, series_acc_niter = 300, series_acc_eps = 1e-10)

dWeibullCount_loglik(x, shape, scale, method = c("series_acc", "series_mat", "conv_direct", "conv_naive", "conv_dePril"), time = 1, na.rm = TRUE, conv_steps = 100, conv_extrap = TRUE, series_terms = 50, series_acc_niter = 300, series_acc_eps = 1e-10, weights = NULL)
evWeibullCount(xmax, shape, scale, method = c("series_acc", "series_mat", "conv_direct", "conv_naive", "conv_dePril"), time = 1, conv_steps = 100, conv_extrap = TRUE, series_terms = 50, series_acc_niter = 300, series_acc_eps = 1e-10)

Arguments

x integer (vector), the desired count values.
shape numeric (length 1), shape parameter of the Weibull count.
scale numeric (length 1), scale parameter of the Weibull count.
method character, one of the available methods, see details.
time double, length of the observation window (defaults to 1).
log logical, if TRUE, the log of the probability will be returned.
conv_steps numeric, number of steps used for the extrapolation.
conv_extrap logical, should Richardson extrapolation be applied?
series_terms numeric, number of terms in the series expansion.
series_acc_niter numeric, number of iterations in the Euler-van Wijngaarden algorithm.
series_acc_eps numeric, tolerance of convergence in the Euler-van Wijngaarden algorithm.
na.rm logical, if TRUE NA’s (produced by taking the log of very small probabilities) will be replaced by the smallest allowed probability; default is TRUE.
weights numeric, vector of weights to apply. If NULL, a vector of one’s will be applied.
xmax unsigned integer, maximum count to be used.

Details

Argument method can be used to specify the desired method, as follows:

"series_mat" - series expansion using matrix techniques,
"series_acc" - Euler-van Wijngaarden accelerated series expansion (default),
"conv_direct" - direct convolution method of section 2,
"conv_naive" - naive convolution described in section 3.1,
"conv_dePril" - dePril convolution described in section 3.2.

The arguments have sensible default values.
Value

for \texttt{dWeibullCount}, a vector of probabilities $P(x(i)), i = 1, \ldots, n$, where $n = \text{length}(x)$.

for \texttt{dWeibullCount_loglik}, a double, the log-likelihood of the count process.

for \texttt{evWeibullCount}, a list with components:

\begin{itemize}
    \item \texttt{ExpectedValue} expected value,
    \item \texttt{Variance} variance.
\end{itemize}

\begin{description}
\item[\texttt{dWeibullgammaCount_mat_Covariates}]
    \textit{Univariate Weibull Count Probability with gamma and covariate heterogeneity}
\end{description}

\textbf{Description}

Univariate Weibull Count Probability with gamma and covariate heterogeneity

\textbf{Usage}

\begin{verbatim}
dWeibullgammaCount_mat_Covariates(x, cc, r, alpha, Xcovar, beta, t = 1,
    logFlag = FALSE, jmax = 100L)
\end{verbatim}

\textbf{Arguments}

\begin{itemize}
    \item \texttt{x, cc, t, logFlag, jmax} TODO keywords internal
    \item \texttt{r} numeric shape of the gamma distribution
    \item \texttt{alpha} numeric rate of the gamma distribution
    \item \texttt{Xcovar} matrix covariates value
    \item \texttt{beta} numeric vector of slopes
\end{itemize}

\begin{description}
\item[\texttt{evCount_conv_bi}]
    \textit{Expected value and variance of renewal count process}
\end{description}

\textbf{Description}

Compute numerically expected values and variances of renewal count processes.
Usage

\[
\text{evCount\_conv\_bi}(x_{\text{max}}, \text{distPars}, \text{dist} = c(\text{"weibull"}, \text{"gamma"}, \text{"gengamma"}, \\
\quad \text{"burr"}), \text{method} = c(\text{"dePril"}, \text{"direct"}, \text{"naive"}), \text{nsteps} = 100, \\
\quad \text{time} = 1, \text{extrap} = \text{TRUE})
\]

\[
\text{evCount\_conv\_user}(x_{\text{max}}, \text{distPars}, \text{extrapolPars}, \text{survR}, \\
\quad \text{method} = c(\text{"dePril"}, \text{"direct"}, \text{"naive"}), \text{nsteps} = 100, \text{time} = 1, \\
\quad \text{extrap} = \text{TRUE})
\]

Arguments

\begin{itemize}
  \item \text{xmax} \hspace{1cm} \text{unsigned integer maximum count to be used.}
  \item \text{distPars} \hspace{1cm} \text{TODO}
  \item \text{dist} \hspace{1cm} \text{TODO}
  \item \text{method} \hspace{1cm} \text{TODO}
  \item \text{nsteps} \hspace{1cm} \text{unsiged integer, number of steps used to compute the integral.}
  \item \text{time} \hspace{1cm} \text{double, time at wich to compute the probabilities. Set to 1 by default.}
  \item \text{extrap} \hspace{1cm} \text{logical, if TRUE, Richardson extrapolation will be applied to improve accuracy.}
  \item \text{extrapolPars} \hspace{1cm} \text{ma::vec of length 2. The extrapolation values.}
  \item \text{survR} \hspace{1cm} \text{function, user supplied survival function; should have signature \text{function(t, distPars)}, \\
    \text{where t is a positive real number (the time where the survival function is evaluated) and distPars is a list of distribution parameters. It should return a double value.}}
\end{itemize}

Details

\text{evCount\_conv\_bi} computes the expected value and variance of renewal count processes for the builtin distirbutions of inter-arrival times.

\text{evCount\_conv\_user} computes the expected value and variance for a user specified distribution of the inter-arrival times.

Value

a named list with components "ExpectedValue" and "Variance".

Examples

\[
\text{pwei\_user <- function(tt, distP) \{}
  \hspace{1cm} \text{alpha <- exp(-log(distP[["scale"]]) / distP[["shape"]])}
  \hspace{1cm} \text{pweibull(q = tt, scale = alpha, shape = distP[["shape"]],}
  \hspace{1cm} \quad \text{lower.tail = FALSE)}
\]

## ev convolution Poisson count
\[
\text{lambda <- 2.56}
\]
\[
\text{beta <- 1}
\]
\[
\text{distPars <- list(scale = lambda, shape = beta)}
\]
evbi <- evCount_conv_bi(20, distPars, dist = "weibull")
evbi <- evCount_conv_user(20, distPars, c(2, 2), pwei_user, "dePril")
c(evbi["ExpectedValue"], lambda)
c(evbi["Variance"], lambda)

c(evu["ExpectedValue"], lambda)
c(evu["Variance"], lambda)

## ev convolution weibull count
lambda <- 2.56
beta <- 1.35
distPars <- list(scale = lambda, shape = beta)
evbi <- evCount_conv_bi(20, distPars, dist = "weibull")
evbi <- evCount_conv_user(20, distPars, c(2.35, 2), pwei_user, "dePril")
x <- 1:20
px <- dCount_conv_bi(x, distPars, "weibull", "dePril",
                      nsteps = 100)
ev <- sum(x * px)
var <- sum(x^2 * px) - ev^2

c(evbi["ExpectedValue"], ev)
c(evbi["Variance"], var)
c(evu["ExpectedValue"], ev)
c(evu["Variance"], var)

---

**Fertility data**

**Description**

Fertility data analysed by Winkelmann(1995). The data comes from the second (1985) wave of German Socio-Economic Panel. The sample is formed by 1,243 women aged 44 or older in 1985. The response variable is the number of children per woman and explanatory variables are described in more details below.

**Usage**

fertility

**Format**

A data frame with 9 variables (5 factors, 4 integers) and 1243 observations:

- children integer; response variable: number of children per woman (integer).
- german factor; is the mother German? (yes or no).
- years_school integer; education measured as years of schooling.
voc_train factor; vocational training? (yes or no)
university factor; university education? (yes or no)
religion factor; mother's religion: Catholic, Protestant, Muslim or Others (reference).
rural factor; rural (yes or no?)
year_birth integer; year of birth (last 2 digits)
age_marriage integer; age at marriage

For further details, see Winlemann(1995).

References


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### football

**Football data**

**Description**


**Usage**

football

**Format**

a data.frame with 6 columns and 1104 observations:

- seasonId integer season identifier (year of the first month of competition).
- gameDate POSIXct game date and time.
- homeTeam,awayTeam character home and away team name.
- homeTeamGoals,awayTeamGoals integer number of goals scored by the home and the away team.

**Details**

The data were collected from [http://www.football-data.co.uk/englandm.php](http://www.football-data.co.uk/englandm.php) and slightly formatted and simplified.
frequency_plot  
*Plot a frequency chart*

**Description**

Plot a frequency chart to compare actual and predicted values.

**Usage**

```r
frequency_plot(count_labels, actual, pred, colours)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>count_labels</td>
<td>character, labels to be used.</td>
</tr>
<tr>
<td>actual</td>
<td>numeric, the observed probabilities for the different count specified in count_labels.</td>
</tr>
<tr>
<td>pred</td>
<td>data.frame of predicted values. Should have the same number of rows as actual and one column per model. The columns’ names will be used as labels for the different models.</td>
</tr>
<tr>
<td>colours</td>
<td>character vector of colour codes with length ncol(pred) + 2.</td>
</tr>
</tbody>
</table>

**Details**

In order to compare actual and fitted values, a barchart plot is created. It is the user’s responsibility to provide the count, observed and fitted values.

---

getParNames  
*Return the names of distribution parameters*

**Description**

Return the names of the parameters of a count distribution.

**Usage**

```r
getParNames(dist, ...)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>dist</td>
<td>character, name of the distribution.</td>
</tr>
<tr>
<td>...</td>
<td>parameters to pass when dist == &quot;custom&quot;.</td>
</tr>
</tbody>
</table>

**Value**

character vector with the names of the distribution parameters.
predict.renewal  Predict method for renewal objects

Description

Compute predictions from renewal objects.

Usage

```r
## S3 method for class 'renewal'
predict(object, newdata = NULL, type = c("response", "prob"), se.fit = FALSE, terms = NULL, na.action = na.pass, time = 1, ...)
```

Arguments

- `object`: Object of class inheriting from "lm"
- `newdata`: An optional data frame in which to look for variables with which to predict. If omitted, the fitted values are used.
- `type`: type of prediction. If equal to "response", give the mean probability associated with the individual covariates. If "prob", give the probability of the observed count.
- `se.fit`: A switch indicating if standard errors are required.
- `terms`: If type = "terms", which terms (default is all terms), a character vector.
- `na.action`: function determining what should be done with missing values in newdata. The default is to predict NA.
- `time`: TODO
- `...`: further arguments passed to or from other methods.

Examples

```r
fn <- system.file("extdata", "McShane_Wei_results_boot.RDS", package = "Countr")
object <- readRDS(fn)
data <- object$data
## old data
predOld.response <- predict(object, type = "response", se.fit = TRUE)
predOld.prob <- predict(object, type = "prob", se.fit = TRUE)

## newData (extracted from old Data)
newData <- head(data)
predNew.response <- predict(object, newdata = newData, type = "response", se.fit = TRUE)
predNew.prob <- predict(object, newdata = newData, type = "prob", se.fit = TRUE)

cbind(head(predOld.response$values),
      head(predNew.response$values))
```
head(predOld.response$se$scale),
head(predOld.response$se$shape),
predNew.response$values,
predNew.response$se$scale,
predNew.response$se$shape)

cbind(head(predOld.prob$values),
head(predOld.prob$se$scale),
head(predOld.prob$se$shape),
predNew.prob$values,
predNew.prob$se$scale,
predNew.prob$se$shape)

renewalCoef Get named vector of coefficients for renewal objects

Description

Get named vector of coefficients for renewal objects.

Usage

renewalCoef(object, ...)

Arguments

object an object, there are methods for several classes, see Details.
...

further arguments to be passed to renewalNames, usually something like target = "weibull".

Details

This is a convenience function for constructing named vector of coefficients for renewal count models. Such vectors are needed, for example, for starting values in the model fitting procedures. The simplest way to get a suitably named vector is to take the coefficients of a fitted model but if the fitting procedure requires initial values, this is seemingly a circular situation.

The overall idea is to take coefficients specified by object and transform them to coefficients suitable for a renewal count model as specified by the arguments "...". The provided methods eliminate the need for tedious manual preparation of such vectors and in the most common cases allow the user to do this in a single line.

The default method extracts the coefficients of object using

c <-coef(object) (an error is raised if this fails). It prepares a named numeric vector with names requested by the arguments in "..." and assigns c to the first length(c) elements of the prepared vector. The net effect is that the coefficients of a model can be initialised from the coefficients of a nested model. For example a Poisson regression model can be used to initialise a Weibull count model. Of course the non-zero shape parameter(s) of the Weibull model need to be set separately.

If object is from class glm, the method is identical to the default method.

If object is from class renewalCoefList, its elements are simply concatenated in one long vector.
Refernce


See Also

renewalNames

---

**renewalCoefList**

`Split a vector using the prefixes of the names for grouping`

**Description**

Split a vector using the prefixes of the names for grouping.

**Usage**

`renewalCoefList(coef)`

**Arguments**

`coef` a named vector

**Details**

The names of the coefficients of renewal regression models are prefixed with the names of the parameters to which they refer. This function splits such vectors into a list with one component for each parameter. For example, for a Weibull renewal regression model this will create a list with components "scale" and "shape".

This is a convenience function allowing users to manipulate the coefficients related to a parameter more easily. `renewalCoef` can convert this list back to a vector.

---

**renewalCount**

`Fit renewal count processes regression models`

**Description**

Fit renewal regression models for count data via maximum likelihood.
Usage

renewalCount(formula, data, subset, na.action, weights, offset,
  dist = c("weibull", "weibullgam", "custom", "gamma", "gengamma"),
  anc = NULL, convPars = NULL, link = NULL, time = 1,
  control = renewal.control(...), customPars = NULL,
  seriesPars = NULL, weimethod = NULL, computeHessian = TRUE,
  standardise = FALSE, standardise_scale = 1, model = TRUE,
  y = TRUE, x = FALSE, ...)

Arguments

formula a formula object. If it is a standard formula object, the left hand side specifies
  the response variable and the right hand sides specifies the regression equation
  for the first parameter of the conditional distribution. formula can also be used
to specify the ancilliary regressions, using the operator ‘|’, see Details.
data, subset, na.action,
  arguments controlling formula processing via model.frame.
weights optional numeric vector of weights.
offset optional numeric vector with an a priori known component to be included in the
  linear predictor of the count model. Currently not used.
dist character, built-in distribution to be used as the inter-arrival time distribution or
  "custom" for a user defined distribution, see Details. Currently the built-in dis-
  tributions are "weibull", "weibullgam", "gamma", "gengamma" (generalized-
  gamma) and "burr".
anc a named list of formulas for ancillary regressions, if any, otherwise NULL. The
  formulas associated with the (exact) parameter names are used. The left-hand
  sides of the formulas in anc are ignored.
convPars a list of convolution parameters arguments with slots nsteps, extrap and convMethod,
  see dCount_conv_bi. If NULL, default parameters will be applied.
link named list of character strings specifying the name of the link functions to be
  used in the regression. If NULL, the canonical link function will be used, i.e, log
  if the parameter is supposed to be positive, identity otherwise.
time numeric, time at which the count is observed; default to unity (1).
control a list of control arguments specified via renewal.control.
customPars list, user inputs if dist = "custom", see details.
seriesPars list, series expansion input parameters with slots terms (number of terms in the
  series expansion), iter (number of iteration in the accelerated series expansion
  algorithm) and eps (tolerance in the accelerated series expansion algorithm).
  Only used if dist = "weibull" and weimethod = c("series_mat","series_acc").
weimethod character, computation method to be used if dist = "weibull" or "weibullgam",
  see dWeibullCount and dWeibullgammaCount.
computeHessian logical, should the hessian (and hence the covariance matrix) be computed nu-
  merically at the fitted values.
standardise logical should the covariates be standardised using standardize::standardize() function.
standardise_scale

numeric the desired scale for the covariates; default to 1

model, y, x  
logicals. If TRUE the corresponding components of the fit (model frame, response, model matrix) are returned.

...  
arguments passed to renewal.control in the default setup.

Details

renewal re-uses design and functionality of the basic R tools for fitting regression model (lm, glm) and is highly inspired by hurdle() and zeroinfl() from package pscl. Package Formula is used to handle formulas.

Argument formula is a formula object. In the simplest case its left-hand side (lhs) designates the response variable and the right-hand side the covariates for the first parameter of the distribution (as reported by getParNames). In this case, covariates for the ancilliary parameters are specified using argument anc.

The ancilliary regressions, can also be specified in argument formula by adding them to the right-hand side, separated by the operator ‘|’. For example Y | shape ~ x + y | z can be used in place of the pair Y ~ x + y and anc = list(shape = ~z). In most cases, the name of the second parameter can be omitted, which for this example gives the equivalent Y ~ x + y | z. The actual rule is that if the parameter is missing from the left-hand side, it is inferred from the default parameter list of the distribution.

As another convenience, if the parameters are to to have the same covariates, it is not necessary to repeat the rhs. For example, Y | shape ~ x + y is equivalent to Y | shape ~ x + y | x + y. Note that this is applied only to parameters listed on the lhs, so Y ~ x + y specifies covariates only for the response variable and not any other parameters.

Distributions for inter-arrival times supported internally by this package can be chosen by setting argument "dist" to a suitable character string. Currently the built-in distributions are "weibull", "weibullgam", "gamma", "gengamma" (generalized-gamma) and "burr".

Users can also provide their own inter-arrival distribution. This is done by setting argument "dist" to "custom", specifying the initial values and giving argument customPars as a list with the following components:

parNames character, the names of the parameters of the distribution. The location parameter should be the first one.

survivalFct function object containing the survival function. It should have signature function(t, distPars) where t is the point where the survival function is evaluated and distPars is the list of the distribution parameters. It should return a double value.

extrapolFct function object computing the extrapolation values (numeric of length 2) from the value of the distribution parameters (in distPars). It should have signature function(distPars) and return a numeric vector of length 2. Only required if the extrapolation is set to TRUE in convPars.

Some checks are done to validate customPars but it is user's responsibility to make sure the the functions have the appropriate signatures.

Note: The Weibull-gamma distribution is an experimental version and should be used with care! It is very sensitive to initial values and there is no guarantee of convergence. It has also been
reparameterized in terms of \((1/r, 1/\alpha, c)\) instead of \((r, \alpha, c)\), where \(r\) and \(\alpha\) are the shape and scale of the gamma distribution and \(c\) is the shape of the Weibull distribution.

\textbf{(2017-08-04)(Georgi) experimental feature:} probability residuals in component 'probResiduals'. I also added type 'prob' to the method for residuals() to extract them.

probResiduals[i] is currently \(1 - \text{Prob}(Y[i] \text{ given the covariates})\). "one minus", so that values close to zero are "good". On its own this is probably not very useful but when comparing two models, if one of them has mostly smaller values than the other, there is some reason to claim that the former is superior. For example (see below), gamModel < poisModel in 3:1

\textbf{Value}

An S3 object of class "renewal", which is a list with components including:

- **coefficients**: values of the fitted coefficients.
- **residuals**: vector of weighted residuals \(\omega * (\text{observed} - \text{fitted})\).
- **fitted.values**: vector of fitted means.
- **optim**: data.frame output of optimx.
- **method**: optimisation algorithm.
- **control**: the control arguments, passed to optimx.
- **start**: starting values, passed to optimx.
- **weights**: weights to apply, if any.
- **n**: number of observations (with weights > 0).
- **iterations**: number of iterations in the optimisation algorithm.
- **execTime**: duration of the optimisation.
- **loglik**: log-likelihood of the fitted model.
- **df.residual**: residuals' degrees of freedom for the fitted model.
- **vcoc**: covariance matrix of all coefficients, computed numerically from the hessian at the fitted coefficients (if \text{computeHessian} is TRUE).
- **dist**: name of the inter-arrival distribution.
- **link**: list, inverse link function corresponding to each parameter in the inter-arrival distribution.
- **converged**: logical, did the optimisation algorithm converge?
- **data**: data used to fit the model.
- **formula**: the original formula.
- **call**: the original function call.
- **anc**: named list of formulas to model regression on ancillary parameters.
- **score_fct**: function to compute the vector of scores defined in Cameron(2013) equation 2.94.
- **convPars**: convolution inputs used.
- **customPars**: named list, user passed distribution inputs, see Details.
- **time**: observed window used, default is 1.0 (see inputs).
- **model**: the full model frame (if \text{model} = TRUE).
- **y**: the response count vector (if \text{y} = TRUE).
- **x**: the model matrix (if \text{x} = TRUE).
## References


## Examples

```r
## Not run:
## may take some time to run depending on your CPU
data(football)
wei = renewalCount(formula = homeTeamGoals ~ 1,
data = football, dist = "weibull", weiMethod = "series_acc",
computeHessian = FALSE, control = renewal.control(trace = 0,
method = "nlminb"))

## End(Not run)
```

`renewalNames`  
*Get names of parameters of renewal regression models*

### Description

Get names of parameters of renewal regression models

### Usage

```r
renewalNames(object, ...)
```

### Arguments

- `object`  
an object.

- `...`  
further arguments.

### Details

`renewalNames` gives the a character vector of names of parameters for renewal regression models. There are two main use scenarios: `renewalNames(object, target = "dist")` and `renewalNames(object, ...)`. In the first scenario `target` can be a count distribution, such as "weibull" or a parameter name, such as shape. In this case `renewalNames` transforms coefficient names of object to those specified by target. In the second scenario the argument list is the same that would be used to call `renewalCount`. In this case `renewalNames` returns the names that would be used by `renewalCount` for the coefficients of the fitted model.
renewal_methods  Methods for renewal objects

Description

Methods for renewal objects.

Usage

## S3 method for class 'renewal'
coef(object, ...)

## S3 method for class 'renewal'
vcov(object, ...)

## S3 method for class 'renewal'
residuals(object, type = c("pearson", "response", "prob"), ...)

## S3 method for class 'renewal'
residuals_plot(object, type = c("pearson", "response", "prob"), ...)

## S3 method for class 'renewal'
fitted(object, ...)

## S3 method for class 'renewal'
confint(object, parm, level = 0.95,
type = c("asymptotic", "boot"), bootType = c("norm", "bca", "basic", "perc"), ...)

## S3 method for class 'renewal'
summary(object, ...)

## S3 method for class 'renewal'
print(x, digits = max(3, getOption("digits") - 3), ...)

## S3 method for class 'summary.renewal'
print(x, digits = max(3, getOption("digits") - 3), width = getOption("width"), ...)

## S3 method for class 'renewal'
model.matrix(object, ...)

## S3 method for class 'renewal'
logLik(object, ...)
## S3 method for class 'renewal'
nobs(object, ...)

## S3 method for class 'renewal'
extractAIC(fit, scale, k = 2, ...)

## S3 method for class 'renewal'
addBootSampleObject(object, ...)

## S3 method for class 'renewal'
df.residual(object, ...)

### Arguments

- **object**: an object from class "renewal".
- **...**: further arguments for methods
- **type, parm, level, bootType, x, digits**: see the corresponding generics and section Details.
- **width**: numeric width length
- **fit, scale, k**: same as in the generic.

### Details

Objects from class "renewal" represent fitted count renewal models and are created by calls to "renewalCount()". There are methods for this class for many of the familiar functions for interacting with fitted models.

### Examples

```r
fn <- system.file("extdata", "McShane_Wei_results_boot.RDS", package = "Countr")
object <- readRDS(fn)
class(object) # "renewal"
coef(object)
vcov(object)

## Pearson residuals: rescaled by sd
head(residuals(object, "pearson"))
## response residuals: not rescaled
head(residuals(object, "response"))

head(fitted(object))

## loglik, nobs, AIC, BIC
cl <- as.numeric(logLik(object)), nobs = nobs(object),
AIC = AIC(object), BIC = BIC(object))

asym <- se.coef(object, , "asymptotic")
boot <- se.coef(object, , "boot")
cbind(asym, boot)
```
## CI for coefficients

asy <- confint(object, type = "asymptotic")
## Commenting out for now, see the note in the code of confint.renewal():
## boot <- confint(object, type = "boot", bootType = "norm")
## list(asy = asym, boot = boot)
summary(object);
print(object);
## see renewal_methods
## see renewal_methods

---

### residuals_plot

**Method to visualise the residuals**

**Description**

A method to visualise the residuals

**Usage**

residuals_plot(object, type, ...)

**Arguments**

- **object**: object returned by one of the count-modeling functions
- **type**: character type of residuals to be used.
- **...**: further arguments for methods.

---

### se.coef

**Extract Standard Errors of Model Coefficients**

**Description**

Extract standard errors of model coefficients from objects returned by count-modeling functions.

**Usage**

se.coef(object, parm, type, ...)

## S3 method for class 'renewal'
se.coef(object, parm, type = c("asymptotic", "boot"),
...)

---
Arguments

- **object**: object returned by one of the count-modeling functions
- **par**: parameter's name or index
- **type**: type of standard error: asymptotic normal standard errors ("asymptotic") or bootstrap ("boot").
- ... further arguments for methods.

Details

The method for class "renewal" extracts standard errors of model coefficients from objects returned by renewal. When bootstrap standard error are requested, the function checks for the bootstrap sample in object. If it is not found, the bootstrap sample is created and a warning is issued. Users can choose between asymptotic normal standard errors (asymptotic) or bootstrap (boot).

Value

- a named numeric vector

Examples

```r
## see examples for renewal_methods
```

---

**surv**

*Wrapper to built in survival functions*

Description

Wrapper to built in survival functions

Usage

```r
surv(t, distPars, dist)
```

Arguments

- **t**: double, time point where the survival is to be evaluated at.
- **distPars**: Rcpp::List with distribution specific slots, see details.
- **dist**: character name of the built-in distribution, see details.
Details

The function wraps all built-in survival distributions. User can choose between the `weibull`, `gamma`, `gengamma` (generalized gamma) and `burr` (Burr type XII distribution). It is the user responsibility to pass the appropriate list of parameters as follows:

**weibull** scale (the scale) and shape (the shape) parameters.

**burr** scale (the scale) and shape1 (the shape1) and shape2 (the shape2) parameters.

**gamma** scale (the scale) and shape (the shape) parameter.

**gengamma** mu (location), sigma (scale) and Q (shape) parameters.

Value

a double giving the value of the survival function at time point t at the parameters’ values.

Examples

```r
tt <- 2.5
## weibull
distP <- list(scale = 1.2, shape = 1.16)
alpha <- exp(-log(distP["scale"])/ distP["shape"])
pweibull(q = tt, scale = alpha, shape = distP["shape"],
        lower.tail = FALSE)
surv(tt, distP, "weibull") ## (almost) same

## gamma
distP <- list(shape = 0.5, rate = 1.0 / 0.7)
pgamma(q = tt, rate = distP["rate"], shape = distP["shape"],
        lower.tail = FALSE)
surv(tt, distP, "gamma") ## (almost) same

## generalized gamma
distP <- list(mu = 0.5, sigma = 0.7, Q = 0.7)
flexsurv::pgengamma(q = tt, mu = distP["mu"],
                    sigma = distP["sigma"],
                    Q = distP["Q"],
                    lower.tail = FALSE)
surv(tt, distP, "gengamma") ## (almost) same
```
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