

# Package ‘eventglm’

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**Type** Package

**Title** Regression Models for Event History Outcomes

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**Description** A user friendly, easy to understand way of doing event history regression for marginal estimands of interest, including the cumulative incidence and the restricted mean survival, using the pseudo observation framework for estimation. For a review of the methodology, see Andersen and Pohar Perme (2010) <[doi:10.1177/0962280209105020](https://doi.org/10.1177/0962280209105020)>. The interface uses the well known formulation of a generalized linear model and allows for features including plotting of residuals, the use of sampling weights, and corrected variance estimation.

**License** GPL-3

**URL** <https://sachsmc.github.io/eventglm/>

**BugReports** <https://github.com/sachsmc/eventglm/issues/>

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---

calc_ipcw_pos	<i>Compute inverse probability of censoring weights pseudo observations</i>
---------------	---

---

### Description

Compute inverse probability of censoring weights pseudo observations

### Usage

```
calc_ipcw_pos(mr, time, causen, type, ipcw.method, Gi)
```

**Arguments**

mr	Model response object returned by <a href="#">Surv</a>
time	Max time
causen	Cause of interest (numeric)
type	Outcome type, "cuminc", "survival", or "rmean"
ipcw.method	"binder" or "hajek"
Gi	vector of estimated censoring probabilities

---

check_mod_cens	<i>Error check censoring model</i>
----------------	------------------------------------

---

**Description**

Censoring model must take the same named arguments as the predefined modules (though they do not all have to be used), and return a vector of pseudo observations.

**Usage**

```
check_mod_cens(model.censoring)
```

**Arguments**

model.censoring	censoring model specification as character or function
-----------------	--

---

colon	<i>Chemotherapy for Stage B/C colon cancer</i>
-------	--

---

**Description**

These are data from one of the first successful trials of adjuvant chemotherapy for colon cancer. Levamisole is a low-toxicity compound previously used to treat worm infestations in animals; 5-FU is a moderately toxic (as these things go) chemotherapy agent. There are only one record per patient for the death outcome (or censoring). This is redistributed from the survival package, with a small modification to include only the death outcome.

**Usage**

```
colon
```

**Format**

A data frame with 929 rows and 17 variables:

**id** id

**study** 1 for all patients

**rx** Treatment - Obs(ervation), Lev(amisole), Lev(amisole)+5-FU

**sex** 1=male

**age** in years

**obstruct** obstruction of colon by tumour

**perfor** perforation of colon

**adhere** adherence to nearby organs

**nodes** number of lymph nodes with detectable cancer

**time** days until death or censoring

**status** censoring status

**differ** differentiation of tumour (1=well, 2=moderate, 3=poor)

**extent** Extent of local spread (1=submucosa, 2=muscle, 3=serosa, 4=contiguous structures)

**surg** time from surgery to registration (0=short, 1=long)

**node4** more than 4 positive lymph nodes

**etype** event type: 1=recurrence,2=death

**event** event indicator: censored, death

**See Also**

[colon](#)

---

confint.pseudoglm

*Confidence Intervals for pseudoglm Model Parameters*

---

**Description**

Computes Wald confidence intervals for one or more parameters in a fitted model. Users can specify the type of variance estimate used, with the default being the robust sandwich variance estimator.

**Usage**

```
## S3 method for class 'pseudoglm'
confint(object, parm, level = 0.95, type = "robust", ...)
```

**Arguments**

object	A fitted model object from <a href="#">cumincglm</a> or <a href="#">rmeanglm</a>
parm	a specification of which parameters are to be given confidence intervals, either a vector of numbers or a vector of names. If missing, all parameters are considered.
level	the confidence level required.
type	The type of variance estimate to use, see <a href="#">vcov.pseudoglm</a>
...	Not used

**Value**

A matrix (or vector) with columns giving lower and upper confidence limits for each parameter. These will be labelled as  $(1-\text{level})/2$  and  $1 - (1-\text{level})/2$  in

**Examples**

```
cumincipcw <- cumincglm(survival::Surv(etime, event) ~ age + sex,
  time = 200, cause = "pcm", link = "identity",
  model.censoring = "independent", data = mgus2)
confint(cumincipcw)
```

---

 cumincglm

*Generalized linear models for cumulative incidence*


---

**Description**

Using pseudo observations for the cumulative incidence, this function then runs a generalized linear model and estimates the parameters representing contrasts in the cumulative incidence at a particular set of times (specified by the `time` argument) across covariate values. The link function can be "identity" for estimating differences in the cumulative incidence, "log" for estimating ratios, and any of the other link functions supported by [quasi](#).

**Usage**

```
cumincglm(
  formula,
  time,
  cause = 1,
  link = "identity",
  model.censoring = "independent",
  formula.censoring = NULL,
  ipcw.method = "binder",
  data,
  survival = FALSE,
  weights,
```

```

subset,
na.action,
offset,
control = list(...),
model = FALSE,
x = TRUE,
y = TRUE,
singular.ok = TRUE,
contrasts = NULL,
...
)

```

### Arguments

formula	A formula specifying the model. The left hand side must be a <a href="#">Surv</a> object specifying a right censored survival or competing risks outcome. The status indicator, normally 0=alive, 1=dead. Other choices are TRUE/FALSE (TRUE = death) or 1/2 (2=death). For competing risks, the event variable will be a factor, whose first level is treated as censoring. The right hand side is the usual linear combination of covariates. If there are multiple time points, the special term "tve(.)" can be used to specify that the effect of the variable inside the parentheses will be time varying. In the output this will be represented as the interaction between the time points and the variable.
time	Numeric vector specifying the times at which the cumulative incidence or survival probability effect estimates are desired.
cause	Numeric or character constant specifying the cause indicator of interest.
link	Link function for the cumulative incidence regression model.
model.censoring	Type of model for the censoring distribution. Options are "stratified", which computes the pseudo-observations stratified on a set of categorical covariates, "aareg" for Aalen's additive hazards model, and "coxph" for Cox's proportional hazards model. With those options, we assume that the time to event and event indicator are conditionally independent of the censoring time, and that the censoring model is correctly specified. If "independent", we assume completely independent censoring, i.e., that the time to event and covariates are independent of the censoring time. the censoring time is independent of the covariates in the model. Can also be a custom function, see <a href="#">Details</a> and the "Extending eventglm" vignette.
formula.censoring	A one sided formula (e.g., $\sim x_1 + x_2$ ) specifying the model for the censoring distribution. If NULL, uses the same mean model as for the outcome.
ipcw.method	Which method to use for calculation of inverse probability of censoring weighted pseudo observations. "binder" the default, uses the number of observations as the denominator, while the "hajek" method uses the sum of the weights as the denominator.
data	Data frame in which all variables of formula can be interpreted.

survival	Set to TRUE to use survival (one minus the cumulative incidence) as the outcome. Not available for competing risks models.
weights	an optional vector of 'prior weights' to be used in the fitting process. Should be NULL or a numeric vector.
subset	an optional vector specifying a subset of observations to be used in the fitting process.
na.action	a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of <a href="#">options</a> , and is <a href="#">na.fail</a> if that is unset. The 'factory-fresh' default is <a href="#">na.omit</a> . Another possible value is NULL, no action. Value <a href="#">na.exclude</a> can be useful.
offset	this can be used to specify an a priori known component to be included in the linear predictor during fitting. This should be NULL or a numeric vector of length equal to the number of cases. One or more <a href="#">offset</a> terms can be included in the formula instead or as well, and if more than one is specified their sum is used. See <a href="#">model.offset</a> . If <code>length(time) &gt; 1</code> , then any offset terms must appear in the formula.
control	a list of parameters for controlling the fitting process. This is passed to <a href="#">glm.control</a> .
model	a logical value indicating whether model frame should be included as a component of the returned value.
x	logical value indicating whether the model matrix used in the fitting process should be returned as components of the returned value.
y	logical value indicating whether the response vector (pseudo-observations) used in the fitting process should be returned as components of the returned value.
singular.ok	logical; if FALSE a singular fit is an error.
contrasts	an optional list. See the contrasts.arg of <a href="#">model.matrix.default</a> .
...	Other arguments passed to <a href="#">glm.fit</a>

## Details

The argument "model.censoring" determines how the pseudo observations are calculated. This can be the name of a function or the function itself, which must have arguments "formula", "time", "cause", "data", "type", "formula.censoring", and "ipcw.method". If it is the name of a function, this code will look for a function with the prefix "pseudo\_" first, to avoid clashes with related methods such as `coxph`. The function then must return a vector of pseudo observations, one for each subject in data which are used in subsequent calculations. For examples of the implementation, see the "pseudo-modules.R" file, or the vignette "Extending eventglm".

## Value

A `pseudoglm` object, with its own methods for `print`, `summary`, and `vcov`. It inherits from `glm`, so `predict` and other `glm` methods are supported.

## Examples

```
cumincipcw <- cumingglm(Surv(etime, event) ~ age + sex,
  time = 200, cause = "pcm", link = "identity",
```

```

      model.censoring = "independent", data = mgus2)
# stratified on only the categorical covariate
  cumincipw2 <- cumincglm(Surv(etime, event) ~ age + sex,
      time = 200, cause = "pcm", link = "identity",
      model.censoring = "stratified",
      formula.censoring = ~ sex, data = mgus2)
# multiple time points
cuminct2 <- cumincglm(Surv(etime, event) ~ age + sex,
  time = c(50, 100, 200), cause = "pcm", link = "identity",
  model.censoring = "independent", data = mgus2)
cuminct3 <- cumincglm(Surv(etime, event) ~ age + tve(sex),
  time = c(50, 100, 200), cause = "pcm", link = "identity",
  model.censoring = "independent", data = mgus2)

```

---

eventglm

*Regression Models for Event History Outcomes*


---

### Description

A user friendly, easy to understand way of doing event history regression for marginal estimands of interest, including the cumulative incidence and the restricted mean survival, using the pseudo observation framework for estimation. The interface uses the well known formulation of a generalized linear model and allows for features including plotting of residuals, the use of sampling weights, and corrected variance estimation.

---

get\_pseudo\_cuminc

*Utility to get jackknife pseudo observations of cumulative incidence*


---

### Description

Utility to get jackknife pseudo observations of cumulative incidence

### Usage

```
get_pseudo_cuminc(marginal.estimate, time, cause, mr)
```

### Arguments

marginal.estimate	
time	A survfit object with no covariates
cause	Time at which to calculate the obs
mr	which cause
	Model response of the survival object

### Value

A vector of pseudo-observations

---

get\_pseudo\_rmean      *Utility to get jackknife pseudo observations of restricted mean*

---

**Description**

Utility to get jackknife pseudo observations of restricted mean

**Usage**

```
get_pseudo_rmean(marginal.estimate, time, cause, mr)
```

**Arguments**

marginal.estimate	A survfit object with no covariates
time	Time at which to calculate the obs
cause	which cause
mr	Model response of the survival object

**Value**

A vector of pseudo-observations

---

jackknife.competing.risks2  
*Compute jackknife pseudo-observations of the cause-specific cumulative incidence for competing risks*

---

**Description**

Compute jackknife pseudo-observations of the cause-specific cumulative incidence for competing risks

**Usage**

```
jackknife.competing.risks2(object, times, cause, mr)
```

**Arguments**

object	A survfit object, with competing events
times	Times at which the cumulative incidence is computed, must be length 1
cause	Value indicating for which cause the cumulative incidence is to be computed, it must match one of the values available in object (see example)
mr	Model response, the result of a call to Surv, or a matrix with two columns: "time" (observed follow up time) and "status" (0 = censored, 1, ..., k = event types)

**Value**

A vector of jackknifed pseudo-observations of the cause-specific cumulative incidence at time times

**Examples**

```
sfit.cuminc <- survival::survfit(survival::Surv(etime, event) ~ 1, data = mgus2)
mrs <- with(mgus2, Surv(etime, event))
pseudo.obs <- jackknife.competing.risks2(sfit.cuminc, times = 200, cause = "pcm", mrs)
mean(pseudo.obs)
# agrees with
summary(sfit.cuminc, times = 200)
```

---

jackknife.survival2    *Compute jackknife pseudo-observations of the survival function*

---

**Description**

Compute jackknife pseudo-observations of the survival function

**Usage**

```
jackknife.survival2(object, times, mr)
```

**Arguments**

object	A survfit object, with a single event (no competing risks)
times	Times at which the survival is computed, must be length 1
mr	Model response, the result of a call to Surv, or a matrix with two columns: "time" (observed follow up time) and "status" (0 = censored, 1 = event)

**Value**

A vector of jackknifed estimates of survival at time times

**Examples**

```
sfit.surv <- survival::survfit(survival::Surv(time, status) ~ 1, data = colon)
mrs <- with(colon, Surv(time, status))
pseudo.obs <- jackknife.survival2(sfit.surv, times = 1000, mrs)
mean(pseudo.obs)
# agrees with
summary(sfit.surv, times = 1000)
```

---

```
leaveOneOut.competing.risks
```

*Compute jackknife pseudo-observations of the cause-specific cumulative incidence for competing risks*

---

### Description

This version computes them for all times up to times, for the restricted mean lifetime lost

### Usage

```
leaveOneOut.competing.risks(object, times, cause, mr)
```

### Arguments

object	A survfit object, with competing events
times	Times at which the cumulative incidence is computed, must be length 1
cause	Value indicating for which cause the cumulative incidence is to be computed, it must match one of the values available in object (see example)
mr	Model response, the result of a call to Surv, or a matrix with two columns: "time" (observed follow up time) and "status" (0 = censored, 1, ..., k = event types)

### Value

A vector of jackknifed values of the cause-specific cumulative incidence at time times

### Examples

```
sfit.cuminc <- survival::survfit(survival::Surv(etime, event) ~ 1, data = mgus2)
mrs <- with(mgus2, Surv(etime, event))
jackvals <- leaveOneOut.competing.risks(sfit.cuminc, times = 200, cause = "pcm", mrs)
```

---

```
leaveOneOut.competing.risks2
```

*Compute jackknife pseudo-observations of the cause-specific cumulative incidence for competing risks*

---

### Description

Compute jackknife pseudo-observations of the cause-specific cumulative incidence for competing risks

### Usage

```
leaveOneOut.competing.risks2(object, times, cause, mr)
```

**Arguments**

object	A survfit object, with competing events
times	Times at which the cumulative incidence is computed, must be length 1
cause	Value indicating for which cause the cumulative incidence is to be computed, it must match one of the values available in object (see example)
mr	Model response, the result of a call to Surv, or a matrix with two columns: "time" (observed follow up time) and "status" (0 = censored, 1, ..., k = event types)

**Value**

A vector of jackknifed values of the cause-specific cumulative incidence at time times

**Examples**

```
sfit.cuminc <- survival::survfit(survival::Surv(etime, event) ~ 1, data = mgus2)
mrs <- with(mgus2, Surv(etime, event))
jackvals <- leaveOneOut.competing.risks2(sfit.cuminc, times = 200, cause = "pcm", mrs)
```

---

leaveOneOut.survival *Compute leave one out jackknife contributions of the survival function*

---

**Description**

For each subject, the survival function is recomputed leaving that subject out. This one does the calculation for all observed times, for calculation of the restricted mean

**Usage**

```
leaveOneOut.survival(object, times, mr)
```

**Arguments**

object	A survfit object, with a single event (no competing risks)
times	Compute values at observed times up to and including this time
mr	Model response, the result of a call to Surv, or a matrix with two columns: "time" (observed follow up time) and "status" (0 = censored, 1 = event)

**Value**

A vector of jackknifed values of survival at time times

**Examples**

```
sfit.surv <- survival::survfit(survival::Surv(time, status) ~ 1, data = colon)
mrs <- with(colon, Surv(time, status))
jackvals <- leaveOneOut.survival(sfit.surv, 1000, mrs)
```

---

leaveOneOut.survival2 *Compute leave one out jackknife contributions of the survival function*

---

### Description

For each subject, the survival function is recomputed leaving that subject out. This is the workhorse for [jackknife.survival2](#) and will generally not be called by the user.

### Usage

```
leaveOneOut.survival2(object, times, mr)
```

### Arguments

object	A survfit object, with a single event (no competing risks)
times	Times at which the survival is computed, must be length 1
mr	Model response, the result of a call to Surv, or a matrix with two columns: "time" (observed follow up time) and "status" (0 = censored, 1 = event)

### Value

A vector of jackknifed values of survival at time times

### Examples

```
sfit.surv <- survival::survfit(survival::Surv(time, status) ~ 1, data = colon)
mrs <- with(colon, Surv(time, status))
jackvals <- leaveOneOut.survival2(sfit.surv, times = 1000, mrs)
```

---

match\_cause *Match cause specification against model response*

---

### Description

Match cause specification against model response

### Usage

```
match_cause(mr, cause)
```

### Arguments

mr	model.response as returned by <a href="#">Surv</a>
cause	Numeric or string indicating the cause of interest

---

`mgus2`*Monoclonal gammopathy data*

---

**Description**

Natural history of 1341 sequential patients with monoclonal gammopathy of undetermined significance (MGUS). This is a superset of the mgus data, at a later point in the accrual process. This dataset is redistributed from the survival package with an added competing risks event indicator.

**Usage**`mgus2`**Format**

A data frame with 1384 observations on the following 10 variables.

`id` subject identifier

`age` age at diagnosis, in years

`sex` a factor with levels F M

`dxyr` year of diagnosis

`hgb` hemoglobin

`creat` creatinine

`mspike` size of the monoclonal serum spike

`ptime` time until progression to a plasma cell malignancy (PCM) or last contact, in months

`pstat` occurrence of PCM: 0=no, 1=yes

`futime` time until death or last contact, in months

`death` occurrence of death: 0=no, 1=yes

`etime` time until either death, pcm, or last contact

`event` factor indicating which event occurred first

**See Also**

[mgus2](#)

---

print.pseudoglm	<i>Print method for pseudoglm</i>
-----------------	-----------------------------------

---

**Description**

Print method for pseudoglm

**Usage**

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

**Arguments**

x	A pseudoglm object, as returned by <a href="#">cumincglm</a> or <a href="#">rmeanglm</a>
digits	Number of significant digits
...	Not used

**Value**

x, invisibly

---

pseudo_aareg	<i>Compute censoring weighted pseudo observations</i>
--------------	---

---

**Description**

Assuming that the censoring depends on covariates, the pseudo observations are calculated with the inverse probability of censoring weighted approach, where the censoring probabilities are estimated using Aalen's additive hazards model.

**Usage**

```
pseudo_aareg(
  formula,
  time,
  cause = 1,
  data,
  type = c("cuminc", "survival", "rmean"),
  formula.censoring = NULL,
  ipcw.method = NULL
)
```

**Arguments**

formula	A formula specifying the outcome model. The left hand side must be a <a href="#">Surv</a> object specifying a right censored survival or competing risks outcome. The status indicator, normally 0=alive, 1=dead. Other choices are TRUE/FALSE (TRUE = death) or 1/2 (2=death). For competing risks, the event variable will be a factor, whose first level is treated as censoring. The right hand side is the usual linear combination of covariates.
time	Numeric constant specifying the time at which the cumulative incidence or survival probability effect estimates are desired.
cause	Numeric or character constant specifying the cause indicator of interest.
data	Data frame in which all variables of formula can be interpreted.
type	One of "survival", "cuminc", or "rmean"
formula.censoring	A right-sided formula specifying which variables to use in the model for the censoring distribution.
ipcw.method	Which method to use for calculation of inverse probability of censoring weighted pseudo observations. "binder" the default, uses the number of observations as the denominator, while the "hajek" method uses the sum of the weights as the denominator.

**Value**

A vector of pseudo observations

**See Also**

[aareg](#)

**Examples**

```
P0i <- pseudo_aareg(Surv(time, status) ~ 1, 1500, cause = 1,
  data = colon, type = "rmean", formula.censoring = ~ sex + age,
  ipcw.method = "binder")

mean(P0i)
```

---

pseudo\_coxph

*Compute censoring weighted pseudo observations*

---

**Description**

Assuming that the censoring depends on covariates, the pseudo observations are calculated with the inverse probability of censoring weighted approach, where the censoring probabilities are estimated using Cox's proportional hazards model.

**Usage**

```

pseudo_coxph(
  formula,
  time,
  cause = 1,
  data,
  type = c("cuminc", "survival", "rmean"),
  formula.censoring = NULL,
  ipcw.method = NULL
)

```

**Arguments**

formula	A formula specifying the outcome model. The left hand side must be a <a href="#">Surv</a> object specifying a right censored survival or competing risks outcome. The status indicator, normally 0=alive, 1=dead. Other choices are TRUE/FALSE (TRUE = death) or 1/2 (2=death). For competing risks, the event variable will be a factor, whose first level is treated as censoring. The right hand side is the usual linear combination of covariates.
time	Numeric constant specifying the time at which the cumulative incidence or survival probability effect estimates are desired.
cause	Numeric or character constant specifying the cause indicator of interest.
data	Data frame in which all variables of formula can be interpreted.
type	One of "survival", "cuminc", or "rmean"
formula.censoring	A right-sided formula specifying which variables to use in the model for the censoring distribution.
ipcw.method	Which method to use for calculation of inverse probability of censoring weighted pseudo observations. "binder" the default, uses the number of observations as the denominator, while the "hajek" method uses the sum of the weights as the denominator.

**Value**

A vector of pseudo observations

**See Also**

[coxph](#)

**Examples**

```

P0i <- pseudo_coxph(Surv(time, status) ~ 1, 1500, cause = 1,
  data = colon, type = "survival", formula.censoring = ~ sex + age,
  ipcw.method = "hajek")

mean(P0i)

```

---

pseudo\_independent      *Compute pseudo observations under independent censoring*

---

### Description

Assuming completely independent censoring, i.e., censoring does not depend on the survival time nor any covariates in the model, the pseudo observations are calculated with the standard jackknife approach

### Usage

```
pseudo_independent(
  formula,
  time,
  cause = 1,
  data,
  type = c("cuminc", "survival", "rmean"),
  formula.censoring = NULL,
  ipcw.method = NULL
)
```

### Arguments

formula	A formula specifying the model. The left hand side must be a <a href="#">Surv</a> object specifying a right censored survival or competing risks outcome. The status indicator, normally 0=alive, 1=dead. Other choices are TRUE/FALSE (TRUE = death) or 1/2 (2=death). For competing risks, the event variable will be a factor, whose first level is treated as censoring. The right hand side is the usual linear combination of covariates.
time	Numeric constant specifying the time at which the cumulative incidence or survival probability effect estimates are desired.
cause	Numeric or character constant specifying the cause indicator of interest.
data	Data frame in which all variables of formula can be interpreted.
type	One of "survival", "cuminc", or "rmean"
formula.censoring	Not used with this method, see <a href="#">pseudo_stratified</a> , <a href="#">pseudo_aareg</a> or <a href="#">pseudo_coxph</a>
ipcw.method	Not used with this method

### Value

A vector of jackknife pseudo observations

### Examples

```
P0i <- pseudo_independent(Surv(time, status) ~ 1, 1500, cause = 1, data = colon, type = "survival")
mean(P0i)
```

---

pseudo\_rmst2                    *Compute pseudo-observations for the restricted mean survival*

---

### Description

Compute pseudo-observations for the restricted mean survival

### Usage

```
pseudo_rmst2(sfit, jacks, times, tmax, type = "cuminc")
```

### Arguments

sfit	A survfit object
jacks	A matrix of leave-one-out jackknife values, subjects in the rows, times in the columns
times	Times at which the survival is calculated
tmax	Max time
type	"cuminc" or "survival"

### Value

A vector of pseudo observations for the restricted mean or lifetime lost

---

pseudo\_stratified                *Compute pseudo observations using stratified jackknife*

---

### Description

Assuming that the censoring depends on covariates with a finite set of levels, the pseudo observations are calculated with the jackknife approach stratified on those covariates.

### Usage

```
pseudo_stratified(
  formula,
  time,
  cause = 1,
  data,
  type = c("cuminc", "survival", "rmean"),
  formula.censoring = NULL,
  ipcw.method = NULL
)
```

**Arguments**

formula	A formula specifying the model. The left hand side must be a <a href="#">Surv</a> object specifying a right censored survival or competing risks outcome. The status indicator, normally 0=alive, 1=dead. Other choices are TRUE/FALSE (TRUE = death) or 1/2 (2=death). For competing risks, the event variable will be a factor, whose first level is treated as censoring. The right hand side is the usual linear combination of covariates.
time	Numeric constant specifying the time at which the cumulative incidence or survival probability effect estimates are desired.
cause	Numeric or character constant specifying the cause indicator of interest.
data	Data frame in which all variables of formula can be interpreted.
type	One of "survival", "cuminc", or "rmean"
formula.censoring	A right-sided formula specifying which variables to stratify on. All variables in this formula must be categorical.
ipcw.method	Not used with this method

**Value**

A vector of jackknife pseudo observations

**Examples**

```
P0i <- pseudo_stratified(Surv(time, status) ~ 1, 1500, cause = 1,
  data = colon, formula.censoring = ~ sex, type = "rmean")
mean(P0i)
```

---

residuals.pseudoglm    *Pseudo-observation scaled residuals*

---

**Description**

Computes residuals according to the recommendations of Pohar-Perme and Andersen (2009) <doi:10.1002/sim.3401>.

**Usage**

```
## S3 method for class 'pseudoglm'
residuals(object, type = NULL, ...)
```

**Arguments**

object	A pseudoglm object, as returned by <a href="#">cumincglm</a> or <a href="#">rmeanglm</a>
type	Either "scaled" (the default for cumulative incidence outcomes) or one of the types available in <a href="#">residuals.glm</a> for restricted mean outcomes, with the default being "deviance".
...	Arguments passed on to <a href="#">residuals.glm</a> .

**Details**

The scaled residuals are computed as

$$\hat{\epsilon}_i = \frac{\hat{E}(V_i) - \hat{Y}_i}{\sqrt{\hat{Y}_i(1 - \hat{Y}_i)}}$$

When the outcome is the cumulative incidence, the denominator corresponds to an estimate of the standard error of the conditional estimate of the outcome in the absence of censoring. For the restricted mean, no such rescaling is done and the computation is passed off to [residuals.glm](#).

**Value**

A numeric vector of residuals

**References**

Perme MP, Andersen PK. Checking hazard regression models using pseudo-observations. *Stat Med*. 2008;27(25):5309-5328. <doi:10.1002/sim.3401>

---

rmeanglm

*Generalized linear models for the restricted mean survival*


---

**Description**

Using pseudo observations for the restricted mean, or the restricted mean lifetime lost in the competing risks case, this function then runs a generalized linear model to estimate associations of the restricted mean/lifetime lost up to a particular time (specified by the `time` argument) with covariates. The link function can be "identity" for estimating differences in the restricted mean, "log" for estimating ratios, and any of the other link functions supported by [quasi](#).

**Usage**

```
rmeanglm(
  formula,
  time,
  cause = 1,
  link = "identity",
  model.censoring = "independent",
  formula.censoring = NULL,
  ipcw.method = "binder",
  data,
  weights,
  subset,
  na.action,
  offset,
  control = list(...),
  model = FALSE,
```

```

x = TRUE,
y = TRUE,
singular.ok = TRUE,
contrasts = NULL,
...
)

```

## Arguments

formula	A formula specifying the model. The left hand side must be a <a href="#">Surv</a> object specifying a right censored survival or competing risks outcome. The status indicator, normally 0=alive, 1=dead. Other choices are TRUE/FALSE (TRUE = death) or 1/2 (2=death). For competing risks, the event variable will be a factor, whose first level is treated as censoring. The right hand side is the usual linear combination of covariates.
time	Numeric constant specifying the time up to which the restricted mean effect estimates are desired.
cause	Numeric or character constant specifying the cause indicator of interest.
link	Link function for the restricted mean regression model.
model.censoring	Type of model for the censoring distribution. Options are "stratified", which computes the pseudo-observations stratified on a set of categorical covariates, "aareg" for Aalen's additive hazards model, and "coxph" for Cox's proportional hazards model. With those options, we assume that the time to event and event indicator are conditionally independent of the censoring time, and that the censoring model is correctly specified. If "independent", we assume completely independent censoring, i.e., that the time to event and covariates are independent of the censoring time. the censoring time is independent of the covariates in the model. Can also be a custom function, see Details and the "Extending eventglm" vignette.
formula.censoring	A one sided formula (e.g., $\sim x_1 + x_2$ ) specifying the model for the censoring distribution. If NULL, uses the same mean model as for the outcome.
ipcw.method	Which method to use for calculation of inverse probability of censoring weighted pseudo observations. "binder" the default, uses the number of observations as the denominator, while the "hajek" method uses the sum of the weights as the denominator.
data	Data frame in which all variables of formula can be interpreted.
weights	an optional vector of 'prior weights' to be used in the fitting process. Should be NULL or a numeric vector.
subset	an optional vector specifying a subset of observations to be used in the fitting process.
na.action	a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of <a href="#">options</a> , and is <a href="#">na.fail</a> if that is unset. The 'factory-fresh' default is <a href="#">na.omit</a> . Another possible value is NULL, no action. Value <a href="#">na.exclude</a> can be useful.

offset	this can be used to specify an a priori known component to be included in the linear predictor during fitting. This should be NULL or a numeric vector of length equal to the number of cases. One or more <code>offset</code> terms can be included in the formula instead or as well, and if more than one is specified their sum is used. See <code>model.offset</code> .
control	a list of parameters for controlling the fitting process. This is passed to <code>glm.control</code> .
model	a logical value indicating whether model frame should be included as a component of the returned value.
x	logical value indicating whether the model matrix used in the fitting process should be returned as components of the returned value.
y	logical value indicating whether the response vector (pseudo-observations) used in the fitting process should be returned as components of the returned value.
singular.ok	logical; if FALSE a singular fit is an error.
contrasts	an optional list. See the <code>contrasts.arg</code> of <code>model.matrix.default</code> .
...	Other arguments passed to <code>glm.fit</code>

## Details

The argument "model.censoring" determines how the pseudo observations are calculated. This can be the name of a function or the function itself, which must have arguments "formula", "time", "cause", "data", "type", "formula.censoring", and "ipcw.method". If it is the name of a function, this code will look for a function with the prefix "pseudo\_" first, to avoid clashes with related methods such as `coxph`. The function then must return a vector of pseudo observations, one for each subject in data which are used in subsequent calculations. For examples of the implementation, see the "pseudo-modules.R" file, or the vignette "Extending eventglm".

## Value

A pseudoglm object, with its own methods for print, summary, and vcov. It inherits from `glm`, so `predict` and other `glm` methods are supported.

## Examples

```
cumincipw <- rmeanglm(Surv(etime, event) ~ age + sex,
  time = 200, cause = "pcm", link = "identity",
  model.censoring = "independent", data = mgus2)
# stratified on only the categorical covariate
cumincipw2 <- rmeanglm(Surv(etime, event) ~ age + sex,
  time = 200, cause = "pcm", link = "identity",
  model.censoring = "stratified",
  formula.censoring = ~ sex, data = mgus2)
```

---

summary.pseudoglm      *Summary method*

---

### Description

Summary method

### Usage

```
## S3 method for class 'pseudoglm'
summary(
  object,
  correlation = FALSE,
  symbolic.cor = FALSE,
  type = "robust",
  ...
)
```

### Arguments

object	A pseudoglm object, as returned by <a href="#">cumincglm</a> or <a href="#">rmeanglm</a>
correlation	logical; if TRUE, the correlation matrix of the estimated parameters is returned and printed.
symbolic.cor	logical; If TRUE, print the correlations in a symbolic form rather than as numbers.
type	The method to use for variance estimation; one of "corrected", "robust", "naive", or "cluster"
...	Not used

### Value

An object of class [summary.glm](#)

---

vcov.pseudoglm      *Compute covariance matrix of regression coefficient estimates*

---

### Description

Compute covariance matrix of regression coefficient estimates

### Usage

```
## S3 method for class 'pseudoglm'
vcov(object, type = "robust", ...)
```

**Arguments**

object	A pseudoglm object, as returned by <a href="#">cumingglm</a> or <a href="#">rmeanglm</a> .
type	The method to use for variance estimation; one of "corrected", "robust", "naive", or "cluster"
...	Not used

**Details**

The "corrected" variance estimate for the cumulative incidence is as described in Overgaard et al. (2017) <doi:10.1214/16-AOS1516>, with code adapted from Overgaard's Stata program. This method does not handle ties and only has marginal benefits in reasonable sample sizes. The default is "robust" which uses the sandwich estimator vcovHC as implemented in the sandwich package. "cluster" is another option if you have clustered observations that uses the vcovCL function in sandwich. Finally "naive" uses the same method as glm to compute the variance, and is known to be anti-conservative. The bootstrap is another recommended option that can be implemented using other tools; there is an example in the vignette.

**Value**

A numeric matrix containing the variance-covariance estimates

**References**

Overgaard, Morten; Parner, Erik Thorlund; Pedersen, Jan. Asymptotic theory of generalized estimating equations based on jack-knife pseudo-observations. *Ann. Statist.* 45 (2017), no. 5, 1988–2015. <doi:10.1214/16-AOS1516>.

**See Also**

[vcovHC](#), [vcovCL](#)

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