Package 'broom'

May 17, 2024

Type Package

Title Convert Statistical Objects into Tidy Tibbles

Version 1.0.6

Description Summarizes key information about statistical objects in tidy tibbles. This makes it easy to report results, create plots and consistently work with large numbers of models at once. Broom provides three verbs that each provide different types of information about a model. tidy() summarizes information about model components such as coefficients of a regression. glance() reports

information about an entire model, such as goodness of fit measures like AIC and BIC. augment() adds information about individual

observations to a dataset, such as fitted values or influence measures.

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URL https://broom.tidymodels.org/, https://github.com/tidymodels/broom

BugReports https://github.com/tidymodels/broom/issues

Depends R (>= 3.5)

- **Imports** backports, dplyr (>= 1.0.0), generics (>= 0.0.2), glue, lifecycle, purrr, rlang, stringr, tibble (>= 3.0.0), tidyr (>= 1.0.0)
- Suggests AER, AUC, bbmle, betareg, biglm, binGroup, boot, btergm (>= 1.10.6), car, carData, caret, cluster, cmprsk, coda, covr, drc, e1071, emmeans, epiR, ergm (>= 3.10.4), fixest (>= 0.9.0), gam (>= 1.15), gee, geepack, ggplot2, glmnet, glmnetUtils, gmm, Hmisc, irlba, interp, joineRML, Kendall, knitr, ks, Lahman, lavaan, leaps, lfe, lm.beta, lme4, lmodel2, lmtest (>= 0.9.38), lsmeans, maps, MASS, mclust, mediation, metafor, mfx, mgcv, mlogit, modeldata, modeltests (>= 0.1.6), muhaz, multcomp, network, nnet, orcutt (>= 2.2), ordinal, plm, poLCA, psych, quantreg, rmarkdown, robust, robustbase, rsample, sandwich, spdep (>= 1.1), spatialreg, speedglm, spelling, survey, survival (>= 3.6-4), systemfit, testthat (>= 2.1.0), tseries, vars, zoo

VignetteBuilder knitr

Config/Needs/website tidyverse/tidytemplate

Encoding UTF-8

RoxygenNote 7.2.3

Language en-US

Collate 'aaa-documentation-helper.R' 'null-and-default-tidiers.R' 'aer-tidiers.R' 'auc-tidiers.R' 'base-tidiers.R' 'bbmle-tidiers.R' 'betareg-tidiers.R' 'biglm-tidiers.R' 'bingroup-tidiers.R' 'boot-tidiers.R' 'broom-package.R' 'broom.R' 'btergm-tidiers.R' 'car-tidiers.R' 'caret-tidiers.R' 'cluster-tidiers.R' 'cmprsk-tidiers.R' 'data-frame-tidiers.R' 'deprecated-0-7-0.R' 'drc-tidiers.R' 'emmeans-tidiers.R' 'epiR-tidiers.R' 'ergm-tidiers.R' 'fixest-tidiers.R' 'gam-tidiers.R' 'geepack-tidiers.R' 'glmnet-cv-glmnet-tidiers.R' 'glmnet-glmnet-tidiers.R' 'gmm-tidiers.R' 'hmisc-tidiers.R' 'joinerml-tidiers.R' 'kendall-tidiers.R' 'ks-tidiers.R' 'lavaan-tidiers.R' 'leaps-tidiers.R' 'lfe-tidiers.R' 'list-irlba.R' 'list-optim-tidiers.R' 'list-svd-tidiers.R' 'list-tidiers.R' 'list-xyz-tidiers.R' 'lm-beta-tidiers.R' 'lmodel2-tidiers.R' 'Imtest-tidiers.R' 'maps-tidiers.R' 'margins-tidiers.R' 'mass-fitdistr-tidiers.R' 'mass-negbin-tidiers.R' 'mass-polr-tidiers.R' 'mass-ridgelm-tidiers.R' 'stats-lm-tidiers.R' 'mass-rlm-tidiers.R' 'mclust-tidiers.R' 'mediation-tidiers.R' 'metafor-tidiers.R' 'mfx-tidiers.R' 'mgcv-tidiers.R' 'mlogit-tidiers.R' 'muhaz-tidiers.R' 'multcomp-tidiers.R' 'nnet-tidiers.R' 'nobs.R' 'orcutt-tidiers.R' 'ordinal-clm-tidiers.R' 'ordinal-clmm-tidiers.R' 'plm-tidiers.R' 'polca-tidiers.R' 'psych-tidiers.R' 'stats-nls-tidiers.R' 'quantreg-nlrq-tidiers.R' 'quantreg-rq-tidiers.R' 'quantreg-rqs-tidiers.R' 'robust-glmrob-tidiers.R' 'robust-lmrob-tidiers.R' 'robustbase-glmrob-tidiers.R' 'robustbase-Imrob-tidiers.R' 'sp-tidiers.R' 'spdep-tidiers.R' 'speedglm-speedglm-tidiers.R' 'speedglm-speedlm-tidiers.R' 'stats-anova-tidiers.R' 'stats-arima-tidiers.R' 'stats-decompose-tidiers.R' 'stats-factanal-tidiers.R' 'stats-glm-tidiers.R' 'stats-htest-tidiers.R' 'stats-kmeans-tidiers.R' 'stats-loess-tidiers.R' 'stats-mlm-tidiers.R' 'stats-prcomp-tidiers.R' 'stats-smooth.spline-tidiers.R' 'stats-summary-Im-tidiers.R' 'stats-time-series-tidiers.R' 'survey-tidiers.R' 'survival-aareg-tidiers.R' 'survival-cch-tidiers.R' 'survival-coxph-tidiers.R' 'survival-pyears-tidiers.R' 'survival-survdiff-tidiers.R' 'survival-survexp-tidiers.R' 'survival-survfit-tidiers.R' 'survival-survreg-tidiers.R' 'systemfit-tidiers.R' 'tseries-tidiers.R' 'utilities.R'

'vars-tidiers.R' 'zoo-tidiers.R' 'zzz.R'

NeedsCompilation no

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augment.betamfx

Augment data with information from a(n) betamfx object

augment.betamfx

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'betamfx'
augment(
    x,
    data = model.frame(x$fit),
    newdata = NULL,
    type.predict = c("response", "link", "precision", "variance", "quantile"),
    type.residuals = c("sweighted2", "deviance", "pearson", "response", "weighted",
        "sweighted"),
    ...
)
```

Arguments

| х | A betamfx object. |
|------|--|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |

| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
|----------------|---|
| type.predict | Character indicating type of prediction to use. Passed to the type argument of <pre>betareg::predict.betareg(). Defaults to "response".</pre> |
| type.residuals | Character indicating type of residuals to use. Passed to the type argument of betareg::residuals.betareg(). Defaults to "sweighted2. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

This augment method wraps augment.betareg() for mfx::betamfx() objects.

Value

A tibble::tibble() with columns:

| .cooksd | Cooks distance. |
|---------|--|
| .fitted | Fitted or predicted value. |
| .resid | The difference between observed and fitted values. |

See Also

```
augment.betareg(), mfx::betamfx()
```

Other mfx tidiers: augment.mfx(), glance.betamfx(), glance.mfx(), tidy.betamfx(), tidy.mfx()

Examples

```
library(mfx)
```

```
# Simulate some data
set.seed(12345)
n <- 1000
x <- rnorm(n)
# Beta outcome
y <- rbeta(n, shape1 = plogis(1 + 0.5 * x), shape2 = (abs(0.2 * x)))
# Use C it be used Yeak it as a set if a s
```

```
# Use Smithson and Verkuilen correction
```

augment.betareg

```
y <- (y * (n - 1) + 0.5) / n
d <- data.frame(y, x)
mod_betamfx <- betamfx(y ~ x | x, data = d)
tidy(mod_betamfx, conf.int = TRUE)
# Compare with the naive model coefficients of the equivalent betareg call (not run)
# tidy(betamfx(y ~ x | x, data = d), conf.int = TRUE)
augment(mod_betamfx)
glance(mod_betamfx)
```

augment.betareg Augment data with information from a(n) betareg object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'betareg'
augment(

```
x,
data = model.frame(x),
newdata = NULL,
type.predict,
type.residuals,
...
)
```

Arguments

| х | A betareg object produced by a call to betareg::betareg(). |
|----------------|--|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| type.predict | Character indicating type of prediction to use. Passed to the type argument of the <pre>stats::predict()</pre> generic. Allowed arguments vary with model class, so be sure to read the predict.my_class documentation. |
| type.residuals | Character indicating type of residuals to use. Passed to the type argument of <pre>stats::residuals()</pre> generic. Allowed arguments vary with model class, so be sure to read the residuals.my_class documentation. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

For additional details on Cook's distance, see stats::cooks.distance().

Value

A tibble::tibble() with columns:

| .cooksd | Cooks distance. |
|---------|--|
| .fitted | Fitted or predicted value. |
| .resid | The difference between observed and fitted values. |

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augment.clm

See Also

augment(), betareg::betareg()

Examples

```
# load libraries for models and data
library(betareg)
# load dats
data("GasolineYield", package = "betareg")
# fit model
mod <- betareg(yield ~ batch + temp, data = GasolineYield)
mod
# summarize model fit with tidiers
tidy(mod)
tidy(mod, conf.int = TRUE)
tidy(mod, conf.int = TRUE, conf.level = .99)
augment(mod)
glance(mod)
```

augment.clm

Augment data with information from a(n) clm object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations. For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'clm'
augment(
    x,
    data = model.frame(x),
    newdata = NULL,
    type.predict = c("prob", "class"),
    ...
)
```

Arguments

| х | A clm object returned from ordinal::clm(). |
|--------------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| type.predict | Which type of prediction to compute, either "prob" or "class", passed to ordinal::predict.clm(). Defaults to "prob". |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

augment.coxph

See Also

```
tidy, ordinal::clm(), ordinal::predict.clm()
```

```
Other ordinal tidiers: augment.polr(), glance.clmm(), glance.clm(), glance.polr(), glance.svyolr(),
tidy.clmm(), tidy.clm(), tidy.polr(), tidy.svyolr()
```

Examples

```
# load libraries for models and data
library(ordinal)
# fit model
fit <- clm(rating ~ temp * contact, data = wine)
# summarize model fit with tidiers
tidy(fit)
tidy(fit, conf.int = TRUE, conf.level = 0.9)
tidy(fit, conf.int = TRUE, conf.type = "Wald", exponentiate = TRUE)
glance(fit)
augment(fit, type.predict = "prob")
augment(fit, type.predict = "class")
# ...and again with another model specification
fit2 <- clm(rating ~ temp, nominal = ~contact, data = wine)
tidy(fit2)
glance(fit2)
```

augment.coxph

Augment data with information from a(n) coxph object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'coxph'
augment(
    x,
    data = model.frame(x),
    newdata = NULL,
    type.predict = "lp",
    type.residuals = "martingale",
    ...
)
```

Arguments

| Х | A coxph object returned from survival::coxph(). |
|----------------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| type.predict | Character indicating type of prediction to use. Passed to the type argument of the stats::predict() generic. Allowed arguments vary with model class, so be sure to read the predict.my_class documentation. |
| type.residuals | Character indicating type of residuals to use. Passed to the type argument of stats::residuals() generic. Allowed arguments vary with model class, so be sure to read the residuals.my_class documentation. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be |

used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

When the modeling was performed with na.action = "na.omit" (as is the typical default), rows with NA in the initial data are omitted entirely from the augmented data frame. When the modeling was performed with na.action = "na.exclude", one should provide the original data as a second argument, at which point the augmented data will contain those rows (typically with NAs in place of the new columns). If the original data is not provided to augment() and na.action = "na.exclude", a warning is raised and the incomplete rows are dropped.

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |
| .se.fit | Standard errors of fitted values. |

See Also

stats::na.action

augment(), survival::coxph()

Other coxph tidiers: glance.coxph(), tidy.coxph()

```
Other survival tidiers: augment.survreg(), glance.aareg(), glance.cch(), glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(), glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(), tidy.survfit(), tidy.survreg()
```

Examples

```
# load libraries for models and data
library(survival)
# fit model
cfit <- coxph(Surv(time, status) ~ age + sex, lung)
# summarize model fit with tidiers
tidy(cfit)
tidy(cfit, exponentiate = TRUE)
lp <- augment(cfit, lung)</pre>
```

```
risks <- augment(cfit, lung, type.predict = "risk")
expected <- augment(cfit, lung, type.predict = "expected")</pre>
glance(cfit)
# also works on clogit models
resp <- levels(logan$occupation)</pre>
n <- nrow(logan)</pre>
indx <- rep(1:n, length(resp))</pre>
logan2 <- data.frame(</pre>
  logan[indx, ],
  id = indx,
  tocc = factor(rep(resp, each = n))
)
logan2$case <- (logan2$occupation == logan2$tocc)</pre>
cl <- clogit(case ~ tocc + tocc:education + strata(id), logan2)</pre>
tidy(cl)
glance(cl)
library(ggplot2)
ggplot(lp, aes(age, .fitted, color = sex)) +
  geom_point()
ggplot(risks, aes(age, .fitted, color = sex)) +
  geom_point()
ggplot(expected, aes(time, .fitted, color = sex)) +
  geom_point()
```

augment.decomposed.ts Augment data with information from a(n) decomposed.ts object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

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Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'decomposed.ts'
augment(x, ...)
```

Arguments

| х | A decomposed.ts object returned from stats::decompose(). |
|---|--|
| | |

... Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in ..., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble with one row for each observation in the original times series:

| .seasonal | The seasonal component of the decomposition. |
|------------|--|
| .trend | The trend component of the decomposition. |
| .remainder | The remainder, or "random" component of the decomposition. |
| .weight | The final robust weights (stl only). |
| .seasadj | The seasonally adjusted (or "deseasonalised") series. |

See Also

augment(), stats::decompose()
Other decompose tidiers: augment.stl()

Examples

```
# time series of temperatures in Nottingham, 1920-1939:
nottem
# perform seasonal decomposition on the data with both decompose
# and stl:
d1 <- decompose(nottem)</pre>
d2 <- stl(nottem, s.window = "periodic", robust = TRUE)</pre>
# compare the original series to its decompositions.
cbind(
  tidy(nottem), augment(d1),
  augment(d2)
)
# visually compare seasonal decompositions in tidy data frames.
library(tibble)
library(dplyr)
library(tidyr)
library(ggplot2)
decomps <- tibble(</pre>
  # turn the ts objects into data frames.
  series = list(as.data.frame(nottem), as.data.frame(nottem)),
  # add the models in, one for each row.
  decomp = c("decompose", "stl"),
  model = list(d1, d2)
) %>%
  rowwise() %>%
  # pull out the fitted data using broom::augment.
  mutate(augment = list(broom::augment(model))) %>%
  ungroup() %>%
  # unnest the data frames into a tidy arrangement of
  # the series next to its seasonal decomposition, grouped
  # by the method (stl or decompose).
  group_by(decomp) %>%
  unnest(c(series, augment)) %>%
  mutate(index = 1:n()) %>%
  ungroup() %>%
  select(decomp, index, x, adjusted = .seasadj)
ggplot(decomps) +
  geom_line(aes(x = index, y = x), colour = "black") +
  geom_line(aes(
   x = index, y = adjusted, colour = decomp,
    group = decomp
  ))
```

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'drc'
augment(
    x,
    data = NULL,
    newdata = NULL,
    se_fit = FALSE,
    conf.int = FALSE,
    conf.level = 0.95,
    ...
```

)

Arguments

```
Х
```

A drc object produced by a call to drc::drm().

| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
|------------|---|
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| se_fit | Logical indicating whether or not a <code>.se.fit</code> column should be added to the augmented output. For some models, this calculation can be somewhat time-consuming. Defaults to FALSE. |
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| .cooksd | Cooks distance. |
|---------|--|
| .fitted | Fitted or predicted value. |
| .lower | Lower bound on interval for fitted values. |
| .resid | The difference between observed and fitted values. |
| .se.fit | Standard errors of fitted values. |
| .upper | Upper bound on interval for fitted values. |

See Also

augment(), drc::drm()

Other drc tidiers: glance.drc(), tidy.drc()

augment.factanal

Examples

```
# load libraries for models and data
library(drc)
# fit model
mod <- drm(dead / total ~ conc, type,
  weights = total, data = selenium, fct = LL.2(), type = "binomial"
)
# summarize model fit with tidiers
tidy(mod)
tidy(mod, conf.int = TRUE)
glance(mod)
augment(mod, selenium)
```

augment.factanal Augment data with information from a(n) factanal object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'factanal'
augment(x, data, ...)
```

Arguments

| х | A factanal object created by stats::factanal(). |
|------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| | Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

When data is not supplied augment.factanal returns one row for each observation, with a factor score column added for each factor X, (.fsX). This is because stats::factanal(), unlike other stats methods like stats::lm(), does not retain the original data.

When data is supplied, augment.factanal returns one row for each observation, with a factor score column added for each factor X, (.fsX).

See Also

augment(), stats::factanal()

Other factanal tidiers: glance.factanal(), tidy.factanal()

augment.felm

Augment data with information from *a*(*n*) felm object

augment.felm

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'felm'
augment(x, data = model.frame(x), ...)

Arguments

| x | A felm object returned from lfe::felm(). |
|------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| | Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • sugment() methods will warn when supplied a newdata argument if it will |

 augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

| A tibble::tibble() with columns: | | |
|----------------------------------|--|--|
| .fitted | Fitted or predicted value. | |
| .resid | The difference between observed and fitted values. | |

See Also

augment(), lfe::felm()
Other felm tidiers: tidy.felm()

Examples

```
# load libraries for models and data
library(lfe)
# use built-in `airquality` dataset
head(airquality)
# no FEs; same as lm()
est0 <- felm(Ozone ~ Temp + Wind + Solar.R, airquality)</pre>
# summarize model fit with tidiers
tidy(est0)
augment(est0)
# add month fixed effects
est1 <- felm(Ozone ~ Temp + Wind + Solar.R | Month, airquality)</pre>
# summarize model fit with tidiers
tidy(est1)
tidy(est1, fe = TRUE)
augment(est1)
glance(est1)
# the "se.type" argument can be used to switch out different standard errors
# types on the fly. In turn, this can be useful exploring the effect of
# different error structures on model inference.
tidy(est1, se.type = "iid")
tidy(est1, se.type = "robust")
# add clustered SEs (also by month)
est2 <- felm(Ozone ~ Temp + Wind + Solar.R | Month | 0 | Month, airquality)</pre>
# summarize model fit with tidiers
tidy(est2, conf.int = TRUE)
tidy(est2, conf.int = TRUE, se.type = "cluster")
tidy(est2, conf.int = TRUE, se.type = "robust")
tidy(est2, conf.int = TRUE, se.type = "iid")
```

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Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'fixest'
augment(
    x,
    data = NULL,
    newdata = NULL,
    type.predict = c("link", "response"),
    type.residuals = c("response", "deviance", "pearson", "working"),
    ...
)
```

Arguments

```
Х
```

A fixest object returned from any of the fixest estimators

| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
|----------------|---|
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| type.predict | Passed to predict.fixest type argument. Defaults to "link" (like predict.glm). |
| type.residuals | Passed to predict.fixest type argument. Defaults to "response" (like residuals.lm, but unlike residuals.glm). |
| | Additional arguments passed to summary and confint. Important arguments are se and cluster. Other arguments are dof, exact_dof, forceCovariance, and keepBounded. See summary.fixest. |

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |

Note

Important note: fixest models do not include a copy of the input data, so you must provide it manually.

augment.fixest only works for fixest::feols(), fixest::feglm(), and fixest::femlm() models. It does not work with results from fixest::fenegbin(), fixest::feNmlm(), or fixest::fepois().

See Also

```
augment(), fixest::feglm(), fixest::femlm(), fixest::feols()
Other fixest tidiers: tidy.fixest()
```

Examples

```
# load libraries for models and data
library(fixest)
gravity <-
   feols(
      log(Euros) ~ log(dist_km) | Origin + Destination + Product + Year, trade
   )
tidy(gravity)</pre>
```

augment.gam

```
glance(gravity)
augment(gravity, trade)
# to get robust or clustered SEs, users can either:
# 1) specify the arguments directly in the `tidy()` call
tidy(gravity, conf.int = TRUE, cluster = c("Product", "Year"))
tidy(gravity, conf.int = TRUE, se = "threeway")
# 2) or, feed tidy() a summary.fixest object that has already accepted
# these arguments
gravity_summ <- summary(gravity, cluster = c("Product", "Year"))
tidy(gravity_summ, conf.int = TRUE)
# approach (1) is preferred.</pre>
```

augment.gam

Augment data with information from *a*(*n*) gam object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'gam'
augment(
    x,
    data = model.frame(x),
    newdata = NULL,
    type.predict,
    type.residuals,
    ...
)
```

Arguments

| х | A gam object returned from a call to mgcv::gam(). |
|----------------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| type.predict | Character indicating type of prediction to use. Passed to the type argument of the <pre>stats::predict()</pre> generic. Allowed arguments vary with model class, so be sure to read the predict.my_class documentation. |
| type.residuals | Character indicating type of residuals to use. Passed to the type argument of <pre>stats::residuals()</pre> generic. Allowed arguments vary with model class, so be sure to read the residuals.my_class documentation. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

For additional details on Cook's distance, see stats::cooks.distance().

Value

A tibble::tibble() with columns:

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augment.glm

| .cooksd | Cooks distance. |
|---------|---|
| .fitted | Fitted or predicted value. |
| .hat | Diagonal of the hat matrix. |
| .resid | The difference between observed and fitted values. |
| .se.fit | Standard errors of fitted values. |
| .sigma | Estimated residual standard deviation when corresponding observation is dropped from model. |

See Also

augment(), mgcv::gam()

Examples

```
# load libraries for models and data
library(mgcv)
# fit model
g <- gam(mpg ~ s(hp) + am + qsec, data = mtcars)
# summarize model fit with tidiers
tidy(g)
tidy(g, parametric = TRUE)
glance(g)
augment(g)</pre>
```

```
augment.glm
```

Augment data with information from a(n) glm object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations. For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'glm'
augment(
    x,
    data = model.frame(x),
    newdata = NULL,
    type.predict = c("link", "response", "terms"),
    type.residuals = c("deviance", "pearson"),
    se_fit = FALSE,
    ...
)
```

Arguments

| х | A glm object returned from stats::glm(). |
|----------------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| type.predict | Passed to stats::predict.glm() type argument. Defaults to "link". |
| type.residuals | Passed to stats::residuals.glm() and to stats::rstandard.glm() type arguments. Defaults to "deviance". |
| se_fit | Logical indicating whether or not a <code>.se.fit</code> column should be added to the augmented output. For some models, this calculation can be somewhat time-consuming. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

If the weights for any of the observations in the model are 0, then columns ".infl" and ".hat" in the result will be 0 for those observations.

A .resid column is not calculated when data is specified via the newdata argument.

Value

A tibble::tibble() with columns:

| .cooksd | Cooks distance. |
|------------|---|
| .fitted | Fitted or predicted value. |
| .hat | Diagonal of the hat matrix. |
| .resid | The difference between observed and fitted values. |
| .se.fit | Standard errors of fitted values. |
| .sigma | Estimated residual standard deviation when corresponding observation is dropped from model. |
| .std.resid | Standardised residuals. |

See Also

stats::glm()

Other Im tidiers: augment.lm(), glance.glm(), glance.lm(), glance.summary.lm(), glance.svyglm(), tidy.glm(), tidy.lm.beta(), tidy.lm(), tidy.mlm(), tidy.summary.lm()

augment.glmRob Augment data with information from a(n) glmRob object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'glmRob'
augment(x, ...)

Arguments

x Unused. ... Unused.

augment.glmrob

Augment data with information from a(n) glmrob object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

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augment.glmrob

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'glmrob'
augment(
    x,
    data = model.frame(x),
    newdata = NULL,
    type.predict = c("link", "response"),
    type.residuals = c("deviance", "pearson"),
    se_fit = FALSE,
    ...
)
```

Arguments

| Х | A glmrob object returned from robustbase::glmrob(). |
|----------------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| type.predict | Character indicating type of prediction to use. Passed to the type argument of the <pre>stats::predict()</pre> generic. Allowed arguments vary with model class, so be sure to read the predict.my_class documentation. |
| type.residuals | Character indicating type of residuals to use. Passed to the type argument of <pre>stats::residuals()</pre> generic. Allowed arguments vary with model class, so be sure to read the residuals.my_class documentation. |
| se_fit | Logical indicating whether or not a <code>.se.fit</code> column should be added to the augmented output. For some models, this calculation can be somewhat time-consuming. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

For tidiers for robust models from the MASS package see tidy.rlm().

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |

See Also

robustbase::glmrob()

Other robustbase tidiers: augment.lmrob(), glance.lmrob(), tidy.glmrob(), tidy.lmrob()

Examples

```
if (requireNamespace("robustbase", quietly = TRUE)) {
 # load libraries for models and data
 library(robustbase)
 data(coleman)
 set.seed(0)
 m <- lmrob(Y ~ ., data = coleman)</pre>
 tidy(m)
 augment(m)
 glance(m)
 data(carrots)
 Rfit <- glmrob(cbind(success, total - success) ~ logdose + block,</pre>
   family = binomial, data = carrots, method = "Mqle",
   control = glmrobMqle.control(tcc = 1.2)
 )
 tidy(Rfit)
 augment(Rfit)
}
```

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'htest'
augment(x, ...)
```

Arguments

>

| х | <pre>An htest objected, such as those created by stats::cor.test(), stats::t.test(), stats::wilcox.test(), stats::chisq.test(), etc.</pre> |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

See stats::chisq.test() for more details on how residuals are computed.

Value

A tibble::tibble() with exactly one row and columns:

| .observed | Observed count. |
|------------|--|
| .prop | Proportion of the total. |
| .row.prop | Row proportion (2 dimensions table only). |
| .col.prop | Column proportion (2 dimensions table only). |
| .expected | Expected count under the null hypothesis. |
| .resid | Pearson residuals. |
| .std.resid | Standardized residual. |
| | |

See Also

augment(), stats::chisq.test()

Other htest tidiers: tidy.htest(), tidy.pairwise.htest(), tidy.power.htest()

Examples

```
tt <- t.test(rnorm(10))
tidy(tt)
# the glance output will be the same for each of the below tests
glance(tt)
tt <- t.test(mpg ~ am, data = mtcars)
tidy(tt)
wt <- wilcox.test(mpg ~ am, data = mtcars, conf.int = TRUE, exact = FALSE)
tidy(wt)
ct <- cor.test(mtcars$wt, mtcars$mpg)
tidy(ct)
chit <- chisq.test(xtabs(Freq ~ Sex + Class, data = as.data.frame(Titanic)))
tidy(chit)
augment(chit)</pre>
```

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'ivreg'
augment(x, data = model.frame(x), newdata = NULL, ...)
```

Arguments

| х | An ivreg object created by a call to AER::ivreg(). |
|---------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

This tidier currently only supports ivreg-classed objects outputted by the AER package. The ivreg package also outputs objects of class ivreg, and will be supported in a later release.

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |

See Also

augment(), AER::ivreg()
Other ivreg tidiers: glance.ivreg(), tidy.ivreg()

Examples

```
# load libraries for models and data
library(AER)
# load data
data("CigarettesSW", package = "AER")
# fit model
ivr <- ivreg(
   log(packs) ~ income | population,
   data = CigarettesSW,
   subset = year == "1995"
)
# summarize model fit with tidiers
tidy(ivr)
tidy(ivr, conf.int = TRUE)
tidy(ivr, conf.int = TRUE, instruments = TRUE)
augment(ivr)
```

. . .

augment.kmeans

```
augment(ivr, data = CigarettesSW)
augment(ivr, newdata = CigarettesSW)
glance(ivr)
```

augment.kmeans

Augment data with information from a(n) kmeans object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'kmeans'
augment(x, data, ...)

Arguments

x A kmeans object created by stats::kmeans().
 data A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data

to the data argument. Augment will report information such as influence and

cooks distance for data passed to the data argument. These measures are only defined for the original training data.

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

.cluster Cluster assignment.

See Also

augment(), stats::kmeans()

Other kmeans tidiers: glance.kmeans(), tidy.kmeans()

Examples

library(cluster)
library(modeldata)
library(dplyr)
data(hpc_data)
x <- hpc_data[, 2:5]
fit <- pam(x, k = 4)
tidy(fit)
glance(fit)
augment(fit, x)</pre>

augment.lm

Augment data with information from *a*(*n*) *lm* object

...

augment.lm

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'lm'
augment(
    x,
    data = model.frame(x),
    newdata = NULL,
    se_fit = FALSE,
    interval = c("none", "confidence", "prediction"),
    conf.level = 0.95,
    ...
)
```

Arguments

х

An lm object created by stats::lm().

data A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. **Do not** pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data.

| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
|------------|--|
| se_fit | Logical indicating whether or not a .se.fit column should be added to the augmented output. For some models, this calculation can be somewhat time-consuming. Defaults to FALSE. |
| interval | Character indicating the type of confidence interval columns to be added to the augmented output. Passed on to predict() and defaults to "none". |
| conf.level | The confidence level to use for the interval created if interval is "confidence" or "prediction". Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence/prediction interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

When the modeling was performed with na.action = "na.omit" (as is the typical default), rows with NA in the initial data are omitted entirely from the augmented data frame. When the modeling was performed with na.action = "na.exclude", one should provide the original data as a second argument, at which point the augmented data will contain those rows (typically with NAs in place of the new columns). If the original data is not provided to augment() and na.action = "na.exclude", a warning is raised and the incomplete rows are dropped.

Some unusual lm objects, such as rlm from MASS, may omit .cooksd and .std.resid. gam from mgcv omits .sigma.

When newdata is supplied, only returns .fitted, .resid and .se.fit columns.

Value

A tibble::tibble() with columns:

| .cooksd | Cooks distance. |
|------------|---|
| .fitted | Fitted or predicted value. |
| .hat | Diagonal of the hat matrix. |
| .lower | Lower bound on interval for fitted values. |
| .resid | The difference between observed and fitted values. |
| .se.fit | Standard errors of fitted values. |
| .sigma | Estimated residual standard deviation when corresponding observation is dropped from model. |
| .std.resid | Standardised residuals. |
| .upper | Upper bound on interval for fitted values. |

augment.lm

See Also

stats::na.action

augment(), stats::predict.lm()

```
Other lm tidiers: augment.glm(), glance.glm(), glance.lm(), glance.summary.lm(), glance.svyglm(), tidy.glm(), tidy.lm(), tidy.mlm(), tidy.summary.lm()
```

Examples

```
library(ggplot2)
library(dplyr)
mod <- lm(mpg ~ wt + qsec, data = mtcars)</pre>
tidy(mod)
glance(mod)
# coefficient plot
d <- tidy(mod, conf.int = TRUE)</pre>
ggplot(d, aes(estimate, term, xmin = conf.low, xmax = conf.high, height = 0)) +
  geom_point() +
  geom_vline(xintercept = 0, lty = 4) +
  geom_errorbarh()
# aside: There are tidy() and glance() methods for lm.summary objects too.
# this can be useful when you want to conserve memory by converting large lm
# objects into their leaner summary.lm equivalents.
s <- summary(mod)</pre>
tidy(s, conf.int = TRUE)
glance(s)
augment(mod)
augment(mod, mtcars, interval = "confidence")
# predict on new data
newdata <- mtcars %>%
  head(6) %>%
  mutate(wt = wt + 1)
augment(mod, newdata = newdata)
# ggplot2 example where we also construct 95% prediction interval
# simpler bivariate model since we're plotting in 2D
mod2 <- lm(mpg ~ wt, data = mtcars)</pre>
au <- augment(mod2, newdata = newdata, interval = "prediction")</pre>
ggplot(au, aes(wt, mpg)) +
  geom_point() +
```

```
geom_line(aes(y = .fitted)) +
 geom_ribbon(aes(ymin = .lower, ymax = .upper), col = NA, alpha = 0.3)
# predict on new data without outcome variable. Output does not include .resid
newdata <- newdata %>%
 select(-mpg)
augment(mod, newdata = newdata)
au <- augment(mod, data = mtcars)</pre>
ggplot(au, aes(.hat, .std.resid)) +
 geom_vline(size = 2, colour = "white", xintercept = 0) +
 geom_hline(size = 2, colour = "white", yintercept = 0) +
 geom_point() +
 geom_smooth(se = FALSE)
plot(mod, which = 6)
ggplot(au, aes(.hat, .cooksd)) +
 geom_vline(xintercept = 0, colour = NA) +
 geom_abline(slope = seq(0, 3, by = 0.5), colour = "white") +
 geom_smooth(se = FALSE) +
 geom_point()
# column-wise models
a <- matrix(rnorm(20), nrow = 10)
b <- a + rnorm(length(a))</pre>
result <- lm(b ~ a)</pre>
tidy(result)
```

augment.lmRob Augment data with information from a(n) lmRob object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

augment.lmRob

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'lmRob'
augment(x, data = model.frame(x), newdata = NULL, ...)

Arguments

| х | A lmRob object returned from robust::lmRob(). |
|---------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| | Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will |
| | |

Details

For tidiers for robust models from the MASS package see tidy.rlm().

be ignored.

See Also

```
robust::lmRob()
Other robust tidiers: glance.glmRob(), glance.lmRob(), tidy.glmRob(), tidy.lmRob()
```

Examples

```
# load modeling library
library(robust)
# fit model
m <- lmRob(mpg ~ wt, data = mtcars)
# summarize model fit with tidiers
tidy(m)
augment(m)
glance(m)</pre>
```

augment.lmrob

Augment data with information from a(n) lmrob object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'lmrob'
augment(x, data = model.frame(x), newdata = NULL, se_fit = FALSE, ...)
```

augment.lmrob

Arguments

| x | A lmrob object returned from robustbase::lmrob(). |
|---------|--|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| se_fit | Logical indicating whether or not a <code>.se.fit</code> column should be added to the augmented output. For some models, this calculation can be somewhat time-consuming. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

For tidiers for robust models from the MASS package see tidy.rlm().

Value

A tibble::tibble() with columns: .fitted Fitted or predicted value.

.resid The difference between observed and fitted values.

See Also

robustbase::lmrob()

Other robustbase tidiers: augment.glmrob(), glance.lmrob(), tidy.glmrob(), tidy.lmrob()

Examples

```
if (requireNamespace("robustbase", quietly = TRUE)) {
    # load libraries for models and data
    library(robustbase)
```

```
data(coleman)
set.seed(0)

m <- lmrob(Y ~ ., data = coleman)
tidy(m)
augment(m)
glance(m)

data(carrots)

Rfit <- glmrob(cbind(success, total - success) ~ logdose + block,
family = binomial, data = carrots, method = "Mqle",
control = glmrobMqle.control(tcc = 1.2)
)

tidy(Rfit)
augment(Rfit)
}</pre>
```

augment.loess Tidy a(n) loess object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'loess'
augment(x, data = model.frame(x), newdata = NULL, se_fit = FALSE, ...)
```

Arguments

| x | A loess objects returned by stats::loess(). |
|---------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |

| se_fit | Logical indicating whether or not a .se.fit column should be added to the augmented output. For some models, this calculation can be somewhat time-consuming. Defaults to FALSE. |
|--------|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

When the modeling was performed with na.action = "na.omit" (as is the typical default), rows with NA in the initial data are omitted entirely from the augmented data frame. When the modeling was performed with na.action = "na.exclude", one should provide the original data as a second argument, at which point the augmented data will contain those rows (typically with NAs in place of the new columns). If the original data is not provided to augment() and na.action = "na.exclude", a warning is raised and the incomplete rows are dropped.

Note that loess objects by default will not predict on data outside of a bounding hypercube defined by the training data unless the original loess object was fit with control = loess.control(surface = \"direct\")). See stats::predict.loess() for details.

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |
| .se.fit | Standard errors of fitted values. |

See Also

stats::na.action

augment(), stats::loess(), stats::predict.loess()

Examples

```
lo <- loess(
  mpg ~ hp + wt,
  mtcars,
  control = loess.control(surface = "direct")
)
augment(lo)</pre>
```

```
# with all columns of original data
augment(lo, mtcars)
# with a new dataset
```

augment(lo, newdata = head(mtcars))

augment.Mclust Augment data with information from a(n) Mclust object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'Mclust'
augment(x, data = NULL, ...)
```

Arguments

```
х
```

An Mclust object return from mclust::Mclust().

| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
|------|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| .class | Predicted class. |
|--------------|---|
| .uncertainty | The uncertainty associated with the classification. Equal to one minus the model class probability. |

See Also

augment(), mclust::Mclust()
Other mclust tidiers: tidy.Mclust()

Examples

load library for models and data
library(mclust)

```
# load data manipulation libraries
library(dplyr)
library(tibble)
library(purrr)
library(tidyr)
set.seed(27)
```

```
centers <- tibble(
  cluster = factor(1:3),
  # number points in each cluster
  num_points = c(100, 150, 50),
  # x1 coordinate of cluster center
  x1 = c(5, 0, -3),
```

```
# x2 coordinate of cluster center
 x^2 = c(-1, 1, -2)
)
points <- centers %>%
 mutate(
    x1 = map2(num_points, x1, rnorm),
    x2 = map2(num_points, x2, rnorm)
 ) %>%
 select(-num_points, -cluster) %>%
 unnest(c(x1, x2))
# fit model
m <- Mclust(points)</pre>
# summarize model fit with tidiers
tidy(m)
augment(m, points)
glance(m)
```

augment.mfx

Augment data with information from *a*(*n*) *mfx* object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

augment.mfx

Usage

```
## S3 method for class 'mfx'
augment(
  х,
  data = model.frame(x$fit),
  newdata = NULL,
  type.predict = c("link", "response", "terms"),
  type.residuals = c("deviance", "pearson"),
  se_fit = FALSE,
  . . .
)
## S3 method for class 'logitmfx'
augment(
 х,
  data = model.frame(x$fit),
  newdata = NULL,
  type.predict = c("link", "response", "terms"),
  type.residuals = c("deviance", "pearson"),
  se_fit = FALSE,
  . . .
)
## S3 method for class 'negbinmfx'
augment(
  х,
  data = model.frame(x$fit),
  newdata = NULL,
  type.predict = c("link", "response", "terms"),
  type.residuals = c("deviance", "pearson"),
  se_fit = FALSE,
  . . .
)
## S3 method for class 'poissonmfx'
augment(
  х,
  data = model.frame(x$fit),
  newdata = NULL,
  type.predict = c("link", "response", "terms"),
  type.residuals = c("deviance", "pearson"),
  se_fit = FALSE,
  . . .
)
## S3 method for class 'probitmfx'
augment(
 х,
```

```
data = model.frame(x$fit),
newdata = NULL,
type.predict = c("link", "response", "terms"),
type.residuals = c("deviance", "pearson"),
se_fit = FALSE,
...
```

Arguments

)

| A logitmfx, negbinmfx, poissonmfx, or probitmfx object. (Note that betamfx objects receive their own set of tidiers.) |
|---|
| A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| Passed to stats::predict.glm() type argument. Defaults to "link". |
| Passed to stats::residuals.glm() and to stats::rstandard.glm() type arguments. Defaults to "deviance". |
| Logical indicating whether or not a .se.fit column should be added to the augmented output. For some models, this calculation can be somewhat time-consuming. Defaults to FALSE. |
| Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| |

Details

This generic augment method wraps augment.glm() for applicable objects from the mfx package.

Value

A tibble::tibble() with columns:

| .cooksd | Cooks distance. |
|---------|----------------------------|
| .fitted | Fitted or predicted value. |

augment.mjoint

| .hat | Diagonal of the hat matrix. |
|------------|---|
| .resid | The difference between observed and fitted values. |
| .se.fit | Standard errors of fitted values. |
| .sigma | Estimated residual standard deviation when corresponding observation is dropped from model. |
| .std.resid | Standardised residuals. |

See Also

```
augment.glm(), mfx::logitmfx(), mfx::negbinmfx(), mfx::poissonmfx(), mfx::probitmfx()
Other mfx tidiers: augment.betamfx(), glance.betamfx(), glance.mfx(), tidy.betamfx(),
tidy.mfx()
```

Examples

```
# load libraries for models and data
library(mfx)
# get the marginal effects from a logit regression
mod_logmfx <- logitmfx(am ~ cyl + hp + wt, atmean = TRUE, data = mtcars)</pre>
tidy(mod_logmfx, conf.int = TRUE)
# compare with the naive model coefficients of the same logit call
tidy(
 glm(am ~ cyl + hp + wt, family = binomial, data = mtcars),
 conf.int = TRUE
)
augment(mod_logmfx)
glance(mod_logmfx)
# another example, this time using probit regression
mod_probmfx <- probitmfx(am ~ cyl + hp + wt, atmean = TRUE, data = mtcars)</pre>
tidy(mod_probmfx, conf.int = TRUE)
augment(mod_probmfx)
glance(mod_probmfx)
```

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'mjoint'
augment(x, data = x\$data, ...)

Arguments

| х | An mjoint object returned from joineRML::mjoint(). |
|------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| | Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will |

 augment() methods will warn when supplied a newdata argument if it will be ignored.

augment.mjoint

Details

See joineRML::fitted.mjoint() and joineRML::residuals.mjoint() for more information on the difference between population-level and individual-level fitted values and residuals.

If fitting a joint model with a single longitudinal process, make sure you are using a named list to define the formula for the fixed and random effects of the longitudinal submodel.

Value

A tibble::tibble() with one row for each original observation with addition columns:

| .fitted_j_0 | population-level fitted values for the j-th longitudinal process |
|-------------|---|
| .fitted_j_1 | individuals-level fitted values for the j-th longitudinal process |
| .resid_j_0 | population-level residuals for the j-th longitudinal process |
| .resid_j_1 | individual-level residuals for the j-th longitudinal process |

Examples

```
# broom only skips running these examples because the example models take a
# while to generatethey should run just fine, though!
## Not run:
```

```
# load libraries for models and data
library(joineRML)
```

```
# fit a joint model with bivariate longitudinal outcomes
data(heart.valve)
```

```
hvd <- heart.valve[!is.na(heart.valve$log.grad) &
!is.na(heart.valve$log.lvmi) &
heart.valve$num <= 50, ]</pre>
```

```
fit <- mjoint(
  formLongFixed = list(
    "grad" = log.grad ~ time + sex + hs,
    "lvmi" = log.lvmi ~ time + sex
  ),
  formLongRandom = list(
    "grad" = ~ 1 | num,
    "lvmi" = ~ time | num
  ),
  formSurv = Surv(fuyrs, status) ~ age,
  data = hvd,
  inits = list("gamma" = c(0.11, 1.51, 0.80)),
  timeVar = "time"
)</pre>
```

extract the survival fixed effects

```
tidy(fit)
# extract the longitudinal fixed effects
tidy(fit, component = "longitudinal")
# extract the survival fixed effects with confidence intervals
tidy(fit, ci = TRUE)
# extract the survival fixed effects with confidence intervals based
# on bootstrapped standard errors
bSE <- bootSE(fit, nboot = 5, safe.boot = TRUE)
tidy(fit, boot_se = bSE, ci = TRUE)
# augment original data with fitted longitudinal values and residuals
hvd2 <- augment(fit)
# extract model statistics
glance(fit)
## End(Not run)</pre>
```

augment.mlogit Augment data with information from a(n) mlogit object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

augment.mlogit

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'mlogit'
augment(x, data = x$model, ...)
```

Arguments

| х | an object returned from mlogit::mlogit(). |
|------|---|
| data | Not currently used |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

At the moment this only works on the estimation dataset. Need to set it up to predict on another dataset.

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|--------------|--|
| .probability | Class probability of modal class. |
| .resid | The difference between observed and fitted values. |

See Also

augment()

Other mlogit tidiers: glance.mlogit(), tidy.mlogit()

Examples

```
# load libraries for models and data
library(mlogit)
data("Fishing", package = "mlogit")
Fish <- dfidx(Fishing, varying = 2:9, shape = "wide", choice = "mode")</pre>
```

```
# fit model
m <- mlogit(mode ~ price + catch | income, data = Fish)
# summarize model fit with tidiers
tidy(m)
augment(m)
glance(m)</pre>
```

augment.nlrq Tidy a(n) nlrq object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'nlrq'
augment(x, data = NULL, newdata = NULL, ...)
```

Arguments

| х | A nlrq object returned from quantreg::nlrq(). |
|---------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

augment.nls

See Also

```
augment(), quantreg::nlrq()
```

```
Other quantreg tidiers: augment.rqs(), augment.rq(), glance.nlrq(), glance.rq(), tidy.nlrq(),
tidy.rqs(), tidy.rq()
```

Examples

```
# fit model
n <- nls(mpg ~ k * e^wt, data = mtcars, start = list(k = 1, e = 2))
# summarize model fit with tidiers + visualization
tidy(n)
augment(n)
glance(n)
library(ggplot2)
ggplot(augment(n), aes(wt, mpg)) +
geom_point() +
geom_line(aes(y = .fitted))
newdata <- head(mtcars)
newdata$wt <- newdata$wt + 1
augment(n, newdata = newdata)
```

augment.nls

Augment data with information from a(n) nls object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations. For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'nls'
augment(x, data = NULL, newdata = NULL, ...)
```

Arguments

| х | An nls object returned from stats::nls(). |
|---------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

augment.nls does not currently support confidence intervals due to a lack of support in stats::predict.nls().

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |

augment.pam

See Also

```
tidy, stats::nls(), stats::predict.nls()
Other nls tidiers: glance.nls(), tidy.nls()
```

Examples

```
# fit model
n <- nls(mpg ~ k * e^wt, data = mtcars, start = list(k = 1, e = 2))
# summarize model fit with tidiers + visualization
tidy(n)
augment(n)
glance(n)
library(ggplot2)
ggplot(augment(n), aes(wt, mpg)) +
geom_point() +
geom_line(aes(y = .fitted))
newdata <- head(mtcars)
newdata $\lefty - newdata$\to t + 1
augment(n, newdata = newdata)
```

augment.pam

Augment data with information from a(n) pam object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations. For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'pam'
augment(x, data = NULL, ...)
```

Arguments

| х | An pam object returned from cluster::pam() |
|------|--|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| .cluster | Cluster assignment. |
|----------|--|
| .fitted | Fitted or predicted value. |
| .resid | The difference between observed and fitted values. |

See Also

augment(), cluster::pam()

Other pam tidiers: glance.pam(), tidy.pam()

augment.plm

Examples

```
# load libraries for models and data
librarv(dplvr)
library(ggplot2)
library(cluster)
library(modeldata)
data(hpc_data)
x <- hpc_data[, 2:5]
p <- pam(x, k = 4)
# summarize model fit with tidiers + visualization
tidy(p)
glance(p)
augment(p, x)
augment(p, x) %>%
  ggplot(aes(compounds, input_fields)) +
  geom_point(aes(color = .cluster)) +
  geom_text(aes(label = cluster), data = tidy(p), size = 10)
```

augment.plm

Augment data with information from a(n) plm object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters

the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'plm'
augment(x, data = model.frame(x), ...)

Arguments

| Х | A plm objected returned by plm::plm(). |
|------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| | Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |

See Also

augment(), plm::plm()

Other plm tidiers: glance.plm(), tidy.plm()

Examples

load libraries for models and data
library(plm)

augment.poLCA

```
# load data
data("Produc", package = "plm")
# fit model
zz <- plm(log(gsp) ~ log(pcap) + log(pc) + log(emp) + unemp,
    data = Produc, index = c("state", "year")
)
# summarize model fit with tidiers
summary(zz)
tidy(zz)
tidy(zz, conf.int = TRUE)
tidy(zz, conf.int = TRUE, conf.level = 0.9)
augment(zz)
glance(zz)
```

augment.poLCA

Augment data with information from a(n) poLCA object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'poLCA'
augment(x, data = NULL, ...)
```

Arguments

| х | A poLCA object returned from poLCA::poLCA(). |
|------|--|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

If the data argument is given, those columns are included in the output (only rows for which predictions could be made). Otherwise, the y element of the poLCA object, which contains the manifest variables used to fit the model, are used, along with any covariates, if present, in x.

Note that while the probability of all the classes (not just the predicted modal class) can be found in the posterior element, these are not included in the augmented output.

Value

A tibble::tibble() with columns:

| .class | Predicted class. |
|--------------|-----------------------------------|
| .probability | Class probability of modal class. |

See Also

augment(), poLCA::poLCA()

Other poLCA tidiers: glance.poLCA(), tidy.poLCA()

augment.poLCA

```
# load libraries for models and data
library(poLCA)
library(dplyr)
# generate data
data(values)
f <- cbind(A, B, C, D) ~ 1
# fit model
M1 <- poLCA(f, values, nclass = 2, verbose = FALSE)
M1
# summarize model fit with tidiers + visualization
tidy(M1)
augment(M1)
glance(M1)
library(ggplot2)
ggplot(tidy(M1), aes(factor(class), estimate, fill = factor(outcome))) +
  geom_bar(stat = "identity", width = 1) +
  facet_wrap(~variable)
# three-class model with a single covariate.
data(election)
f2a <- cbind(
  MORALG, CARESG, KNOWG, LEADG, DISHONG, INTELG,
  MORALB, CARESB, KNOWB, LEADB, DISHONB, INTELB
) ~ PARTY
nes2a <- poLCA(f2a, election, nclass = 3, nrep = 5, verbose = FALSE)</pre>
td <- tidy(nes2a)</pre>
td
ggplot(td, aes(outcome, estimate, color = factor(class), group = class)) +
  geom_line() +
  facet_wrap(~variable, nrow = 2) +
  theme(axis.text.x = element_text(angle = 90, hjust = 1))
au <- augment(nes2a)</pre>
au
count(au, .class)
```

```
# if the original data is provided, it leads to NAs in new columns
# for rows that weren't predicted
au2 <- augment(nes2a, data = election)
au2
dim(au2)
```

augment.polr

Augment data with information from a(n) polr object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'polr'
augment(
    x,
    data = model.frame(x),
    newdata = NULL,
    type.predict = c("class"),
    ...
)
```

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augment.polr

Arguments

| х | A polr object returned from MASS::polr(). |
|---|---|
| A base::data.frame or tibble::tibble() containing the original data the used to produce the object x. Defaults to stats::model.frame(x) augment(my_fit) returns the augmented original data. Do not pass ne to the data argument. Augment will report information such as influen cooks distance for data passed to the data argument. These measures and defined for the original training data. | |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| type.predict | Which type of prediction to compute, passed to MASS:::predict.polr(). Only supports "class" at the moment. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

See Also

tidy(), MASS::polr()

```
Other ordinal tidiers: augment.clm(), glance.clm(), glance.clm(), glance.polr(), glance.svyolr(),
tidy.clm(), tidy.clm(), tidy.polr(), tidy.svyolr()
```

```
# load libraries for models and data
library(MASS)
# fit model
fit <- polr(Sat ~ Infl + Type + Cont, weights = Freq, data = housing)
# summarize model fit with tidiers
tidy(fit, exponentiate = TRUE, conf.int = TRUE)
glance(fit)
augment(fit, type.predict = "class")
fit2 <- polr(factor(gear) ~ am + mpg + qsec, data = mtcars)
tidy(fit, p.values = TRUE)
```

augment.prcomp

Augment data with information from *a*(*n*) prcomp object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'prcomp'
augment(x, data = NULL, newdata, ...)

Arguments

x A prcomp object returned by stats::prcomp().

data A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. **Do not** pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data.

| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
|---------|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| Value | |

A tibble::tibble containing the original data along with additional columns containing each observation's projection into PCA space.

See Also

stats::prcomp(), svd_tidiers

Other svd tidiers: tidy.prcomp(), tidy_irlba(), tidy_svd()

augment.rlm

Augment data with information from a(n) rlm object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'rlm'
augment(x, data = model.frame(x), newdata = NULL, se_fit = FALSE, ...)
```

Arguments

| х | An rlm object returned by MASS::rlm(). |
|---------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| se_fit | Logical indicating whether or not a .se.fit column should be added to the augmented output. For some models, this calculation can be somewhat time-consuming. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|---|
| .hat | Diagonal of the hat matrix. |
| .resid | The difference between observed and fitted values. |
| .se.fit | Standard errors of fitted values. |
| .sigma | Estimated residual standard deviation when corresponding observation is dropped from model. |

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augment.rma

See Also

MASS::rlm()

Other rlm tidiers: glance.rlm(), tidy.rlm()

Examples

```
# load libraries for models and data
library(MASS)
# fit model
r <- rlm(stack.loss ~ ., stackloss)
# summarize model fit with tidiers
tidy(r)
augment(r)
glance(r)
```

augment.rma

Augment data with information from a(n) rma object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'rma'
augment(x, interval = c("prediction", "confidence"), ...)
```

Arguments

| X | An rma object such as those created by metafor::rma(), metafor::rma.uni(), metafor::rma.glmm(), metafor::rma.mh(), metafor::rma.mv(), or metafor::rma.peto(). |
|----------|--|
| interval | For rma.mv models, should prediction intervals ("prediction", default) or con- fidence intervals ("confidence") intervals be returned? For rma.uni models, prediction intervals are always returned. For rma.mh and rma.peto models, confidence intervals are always returned. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|-----------------|---|
| .lower | Lower bound on interval for fitted values. |
| .moderator | In meta-analysis, the moderators used to calculate the predicted values. |
| .moderator.leve | 21 |
| | In meta-analysis, the level of the moderators used to calculate the predicted values. |
| .resid | The difference between observed and fitted values. |
| .se.fit | Standard errors of fitted values. |
| .upper | Upper bound on interval for fitted values. |
| .observed | The observed values for the individual studies |

```
# load modeling library
library(metafor)
# generate data and fit
df <-</pre>
```

augment.rq

```
escalc(
    measure = "RR",
    ai = tpos,
    bi = tneg,
    ci = cpos,
    di = cneg,
    data = dat.bcg
)
meta_analysis <- rma(yi, vi, data = df, method = "EB")
# summarize model fit with tidiers
augment(meta_analysis)</pre>
```

augment.rq

Augment data with information from a(n) rq object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'rq'
augment(x, data = model.frame(x), newdata = NULL, ...)
```

Arguments

| Х | An rq object returned from quantreg::rq(). |
|---------|--|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| | Arguments passed on to quantreg::predict.rq |
| | object object of class rq or rqs or rq.process produced by rq |
| | interval type of interval desired: default is 'none', when set to 'confidence' the function returns a matrix predictions with point predictions for each of the 'newdata' points as well as lower and upper confidence limits. |
| | level converage probability for the 'confidence' intervals. |
| | type For predict.rq, the method for 'confidence' intervals, if desired. If 'per- centile' then one of the bootstrap methods is used to generate percentile intervals for each prediction, if 'direct' then a version of the Portnoy and Zhou (1998) method is used, and otherwise an estimated covariance ma- trix for the parameter estimates is used. Further arguments to determine the choice of bootstrap method or covariance matrix estimate can be passed via the argument. For predict.rqs and predict.rq.process when stepfun = TRUE, type is "Qhat", "Fhat" or "fhat" depending on whether the user would like to have estimates of the conditional quantile, distribution or density functions respectively. As noted below the two former estimates can be monotonized with the function rearrange. When the "fhat" option is invoked, a list of conditional density functions is returned based on Sil- verman's adaptive kernel method as implemented in akj and approxfun. |
| | na.action function determining what should be done with missing values in 'newdata'. The default is to predict 'NA'. |

Details

Depending on the arguments passed on to predict.rq via ..., a confidence interval is also calculated on the fitted values resulting in columns .lower and .upper. Does not provide confidence intervals when data is specified via the newdata argument.

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |
| tau | Quantile. |

augment.rqs

See Also

```
augment, quantreg::rq(), quantreg::predict.rq()
Other quantreg tidiers: augment.nlrq(), augment.rqs(), glance.nlrq(), glance.rq(), tidy.nlrq(),
tidy.rqs(), tidy.rq()
```

Examples

```
# load modeling library and data
library(quantreg)
data(stackloss)
# median (11) regression fit for the stackloss data.
mod1 <- rq(stack.loss ~ stack.x, .5)</pre>
# weighted sample median
mod2 <- rq(rnorm(50) ~ 1, weights = runif(50))</pre>
# summarize model fit with tidiers
tidy(mod1)
glance(mod1)
augment(mod1)
tidy(mod2)
glance(mod2)
augment(mod2)
# varying tau to generate an rqs object
mod3 <- rq(stack.loss ~ stack.x, tau = c(.25, .5))</pre>
tidy(mod3)
augment(mod3)
# glance cannot handle rqs objects like `mod3`--use a purrr
# `map`-based workflow instead
```

augment.rqs

Augment data with information from *a*(*n*) rqs object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'rqs'
augment(x, data = model.frame(x), newdata, ...)
```

Arguments

| х | An rqs object returned from quantreg::rq(). |
|---------|--|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| | Arguments passed on to quantreg::predict.rq |
| | object object of class rq or rqs or rq.process produced by rq interval type of interval desired: default is 'none', when set to 'confidence' the function returns a matrix predictions with point predictions for each of the 'newdata' points as well as lower and upper confidence limits. |
| | level converage probability for the 'confidence' intervals. |
| | type For predict.rq, the method for 'confidence' intervals, if desired. If 'per- centile' then one of the bootstrap methods is used to generate percentile intervals for each prediction, if 'direct' then a version of the Portnoy and |

augment.rqs

Zhou (1998) method is used, and otherwise an estimated covariance matrix for the parameter estimates is used. Further arguments to determine the choice of bootstrap method or covariance matrix estimate can be passed via the ... argument. For predict.rqs and predict.rq.process when stepfun = TRUE, type is "Qhat", "Fhat" or "fhat" depending on whether the user would like to have estimates of the conditional quantile, distribution or density functions respectively. As noted below the two former estimates can be monotonized with the function rearrange. When the "fhat" option is invoked, a list of conditional density functions is returned based on Silverman's adaptive kernel method as implemented in akj and approxfun.

na.action function determining what should be done with missing values in 'newdata'. The default is to predict 'NA'.

Details

Depending on the arguments passed on to predict.rq via ..., a confidence interval is also calculated on the fitted values resulting in columns .lower and .upper. Does not provide confidence intervals when data is specified via the newdata argument.

See Also

```
augment, quantreg::rq(), quantreg::predict.rqs()
```

```
Other quantreg tidiers: augment.nlrq(), augment.rq(), glance.nlrq(), glance.rq(), tidy.nlrq(),
tidy.rqs(), tidy.rq()
```

```
# load modeling library and data
library(quantreg)
data(stackloss)
# median (11) regression fit for the stackloss data.
mod1 <- rq(stack.loss ~ stack.x, .5)</pre>
# weighted sample median
mod2 <- rq(rnorm(50) \sim 1, weights = runif(50))
# summarize model fit with tidiers
tidy(mod1)
glance(mod1)
augment(mod1)
tidy(mod2)
glance(mod2)
augment(mod2)
# varying tau to generate an rqs object
mod3 <- rq(stack.loss ~ stack.x, tau = c(.25, .5))</pre>
```

```
tidy(mod3)
augment(mod3)
# glance cannot handle rqs objects like `mod3`--use a purrr
# `map`-based workflow instead
```

augment.sarlm Augment data with information from a(n) spatialreg object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'sarlm'
augment(x, data = x\$X, ...)

Arguments

| х | An object returned from spatialreg::lagsarlm() or spatialreg::errorsarlm() |
|------|--|
| data | Ignored, but included for internal consistency. See the details below. |

augment.sarlm

. . .

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

The predict method for sarlm objects assumes that the response is known. See ?predict.sarlm for more discussion. As a result, since the original data can be recovered from the fit object, this method currently does not take in data or newdata arguments.

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |

See Also

augment()

Other spatialreg tidiers: glance.sarlm(), tidy.sarlm()

```
# load libraries for models and data
library(spatialreg)
library(spdep)
# load data
data(oldcol, package = "spdep")
listw <- nb2listw(COL.nb, style = "W")
# fit model
crime_sar <-
lagsarlm(CRIME ~ INC + HOVAL,
    data = COL.OLD,
    listw = listw,
    method = "eigen"
)
```

```
# summarize model fit with tidiers
tidy(crime_sar)
tidy(crime_sar, conf.int = TRUE)
glance(crime_sar)
augment(crime_sar)
# fit another model
crime_sem <- errorsarlm(CRIME ~ INC + HOVAL, data = COL.OLD, listw)</pre>
# summarize model fit with tidiers
tidy(crime_sem)
tidy(crime_sem, conf.int = TRUE)
glance(crime_sem)
augment(crime_sem)
# fit another model
crime_sac <- sacsarlm(CRIME ~ INC + HOVAL, data = COL.OLD, listw)</pre>
# summarize model fit with tidiers
tidy(crime_sac)
tidy(crime_sac, conf.int = TRUE)
glance(crime_sac)
augment(crime_sac)
```

augment.smooth.spline Tidy a(n) smooth.spline object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'smooth.spline'
augment(x, data = x$data, ...)
```

Arguments

x A smooth.spline object returned from stats::smooth.spline().

data A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. **Do not** pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data.

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. . .

- Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:
 - tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
 - augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |

See Also

augment(), stats::smooth.spline(), stats::predict.smooth.spline()

Other smoothing spline tidiers: glance.smooth.spline()

```
# fit model
spl <- smooth.spline(mtcars$wt, mtcars$mpg, df = 4)
# summarize model fit with tidiers
augment(spl, mtcars)
# calls original columns x and y
augment(spl)
library(ggplot2)
ggplot(augment(spl, mtcars), aes(wt, mpg)) +
geom_point() +
geom_line(aes(y = .fitted))</pre>
```

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'speedlm'
augment(x, data = model.frame(x), newdata = NULL, ...)

Arguments

| х | A speedlm object returned from speedglm::speedlm(). |
|---------|---|
| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |

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augment.stl

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |

See Also

speedglm::speedlm()

Other speedlm tidiers: glance.speedglm(), glance.speedlm(), tidy.speedglm(), tidy.speedlm()

Examples

```
# load modeling library
library(speedglm)
# fit model
mod <- speedlm(mpg ~ wt + qsec, data = mtcars, fitted = TRUE)
# summarize model fit with tidiers
tidy(mod)
glance(mod)
augment(mod)
```

augment.stl Augment data with information from a(n) stl object

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

S3 method for class 'stl'
augment(x, data = NULL, weights = TRUE, ...)

Arguments

| х | An stl object returned from stats::stl(). |
|---------|---|
| data | Ignored, included for consistency with the augment generic signature only. |
| weights | Logical indicating whether or not to include the robust weights in the output. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble with one row for each observation in the original times series:

| .seasonal | The seasonal component of the decomposition. |
|------------|--|
| .trend | The trend component of the decomposition. |
| .remainder | The remainder, or "random" component of the decomposition. |
| .weight | The final robust weights, if requested. |
| .seasadj | The seasonally adjusted (or "deseasonalised") series. |

See Also

augment(), stats::stl()

Other decompose tidiers: augment.decomposed.ts()

Description

Augment accepts a model object and a dataset and adds information about each observation in the dataset. Most commonly, this includes predicted values in the .fitted column, residuals in the .resid column, and standard errors for the fitted values in a .se.fit column. New columns always begin with a . prefix to avoid overwriting columns in the original dataset.

Users may pass data to augment via either the data argument or the newdata argument. If the user passes data to the data argument, it **must** be exactly the data that was used to fit the model object. Pass datasets to newdata to augment data that was not used during model fitting. This still requires that at least all predictor variable columns used to fit the model are present. If the original outcome variable used to fit the model is not included in newdata, then no .resid column will be included in the output.

Augment will often behave differently depending on whether data or newdata is given. This is because there is often information associated with training observations (such as influences or related) measures that is not meaningfully defined for new observations.

For convenience, many augment methods provide default data arguments, so that augment(fit) will return the augmented training data. In these cases, augment tries to reconstruct the original data based on the model object with varying degrees of success.

The augmented dataset is always returned as a tibble::tibble with the **same number of rows** as the passed dataset. This means that the passed data must be coercible to a tibble. If a predictor enters the model as part of a matrix of covariates, such as when the model formula uses splines::ns(), stats::poly(), or survival::Surv(), it is represented as a matrix column.

We are in the process of defining behaviors for models fit with various na.action arguments, but make no guarantees about behavior when data is missing at this time.

Usage

```
## S3 method for class 'survreg'
augment(
    x,
    data = model.frame(x),
    newdata = NULL,
    type.predict = "response",
    type.residuals = "response",
    ...
)
```

Arguments

```
Х
```

An survreg object returned from survival::survreg().

| data | A base::data.frame or tibble::tibble() containing the original data that was used to produce the object x. Defaults to stats::model.frame(x) so that augment(my_fit) returns the augmented original data. Do not pass new data to the data argument. Augment will report information such as influence and cooks distance for data passed to the data argument. These measures are only defined for the original training data. |
|----------------|--|
| newdata | A base::data.frame() or tibble::tibble() containing all the original pre- dictors used to create x. Defaults to NULL, indicating that nothing has been passed to newdata. If newdata is specified, the data argument will be ignored. |
| type.predict | Character indicating type of prediction to use. Passed to the type argument of the stats::predict() generic. Allowed arguments vary with model class, so be sure to read the predict.my_class documentation. |
| type.residuals | Character indicating type of residuals to use. Passed to the type argument of <pre>stats::residuals()</pre> generic. Allowed arguments vary with model class, so be sure to read the residuals.my_class documentation. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| .fitted | Fitted or predicted value. |
|---------|--|
| .resid | The difference between observed and fitted values. |
| .se.fit | Standard errors of fitted values. |

See Also

augment(), survival::survreg()

Other survreg tidiers: glance.survreg(), tidy.survreg()

Other survival tidiers: augment.coxph(), glance.aareg(), glance.cch(), glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(), glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(), tidy.survfit(), tidy.survreg()

Examples

load libraries for models and data

augment_columns

```
library(survival)
# fit model
sr <- survreg(</pre>
  Surv(futime, fustat) ~ ecog.ps + rx,
  ovarian,
  dist = "exponential"
)
# summarize model fit with tidiers + visualization
tidy(sr)
augment(sr, ovarian)
glance(sr)
# coefficient plot
td <- tidy(sr, conf.int = TRUE)</pre>
library(ggplot2)
ggplot(td, aes(estimate, term)) +
  geom_point() +
  geom_errorbarh(aes(xmin = conf.low, xmax = conf.high), height = 0) +
  geom_vline(xintercept = 0)
```

| augment_columns Add fitted values, residuals, and other common outputs to an augment call | ent |
|---|-----|
|---|-----|

Description

augment_columns is intended for use in the internals of augment methods only and is exported for developers extending the broom package. Please instead use augment() to appropriately make use of the functionality in augment_columns().

Usage

```
augment_columns(
    x,
    data,
    newdata = NULL,
    type,
    type.predict = type,
    type.residuals = type,
    se.fit = TRUE,
    ...
)
```

Arguments

| х | a model |
|----------------|---|
| data | original data onto which columns should be added |
| newdata | new data to predict on, optional |
| type | Type of prediction and residuals to compute |
| type.predict | Type of prediction to compute; by default same as type |
| type.residuals | Type of residuals to compute; by default same as type |
| se.fit | Value to pass to predict's se.fit, or NULL for no value |
| | extra arguments (not used) |

Details

Note that, in the case that a residuals() or influence() generic is not implemented for the supplied model x, the function will fail quietly.

bootstrap

Set up bootstrap replicates of a dplyr operation

Description

The bootstrap() function is deprecated and will be removed from an upcoming release of broom. For tidy resampling, please use the rsample package instead. Functionality is no longer supported for this method.

Usage

bootstrap(df, m, by_group = FALSE)

Arguments

| df | a data frame |
|----------|--|
| m | number of bootstrap replicates to perform |
| by_group | If TRUE, then bootstrap within each group if df is a grouped tibble. |

Details

This code originates from Hadley Wickham (with a few small corrections) here: https://github.com/tidyverse/dplyr/issues/269

See Also

Other deprecated: confint_tidy(), data.frame_tidiers, finish_glance(), fix_data_frame(), summary_tidiers, tidy.density(), tidy.dist(), tidy.ftable(), tidy.numeric() confint_tidy

Description

This function is now deprecated and will be removed from a future release of broom.

Usage

```
confint_tidy(x, conf.level = 0.95, func = stats::confint, ...)
```

Arguments

| х | a model object for which confint() can be calculated |
|------------|--|
| conf.level | confidence level |
| func | A function to compute a confidence interval for x. Calling func(x, level = $conf.level,$) must return an object coercible to a tibble. This dataframe like object should have to columns corresponding the lower and upper bounds on the confidence interval. |
| | extra arguments passed on to confint |

Details

Return a confidence interval as a tidy data frame. This directly wraps the confint() function, but ensures it follows broom conventions: column names of conf.low and conf.high, and no row names.

confint_tidy

Value

A tibble with two columns: conf.low and conf.high.

See Also

```
Other deprecated: bootstrap(), data.frame_tidiers, finish_glance(), fix_data_frame(),
summary_tidiers, tidy.density(), tidy.dist(), tidy.ftable(), tidy.numeric()
```

Description

Data frame tidiers are deprecated and will be removed from an upcoming release of broom.

Usage

```
## S3 method for class 'data.frame'
tidy(x, ..., na.rm = TRUE, trim = 0.1)
## S3 method for class 'data.frame'
augment(x, data, ...)
## S3 method for class 'data.frame'
glance(x, ...)
```

Arguments

| х | A data.frame |
|-------|---|
| | Additional arguments for other methods. |
| na.rm | a logical value indicating whether NA values should be stripped before the com- putation proceeds. |
| trim | the fraction (0 to 0.5) of observations to be trimmed from each end of x before the mean is computed. Passed to the trim argument of mean |
| data | data, not used |

Details

These perform tidy summaries of data.frame objects. tidy produces summary statistics about each column, while glance simply reports the number of rows and columns. Note that augment.data.frame will throw an error.

Value

tidy.data.frame produces a data frame with one row per original column, containing summary statistics of each:

| column | name of original column |
|---------|--|
| n | Number of valid (non-NA) values |
| mean | mean |
| sd | standard deviation |
| median | median |
| trimmed | trimmed mean, with trim defaulting to .1 |

data.frame_tidiers

| mad | median absolute deviation (from the median) | |
|--|---|--|
| min | minimum value | |
| max | maximum value | |
| range | range | |
| skew | skew | |
| kurtosis | kurtosis | |
| se | standard error | |
| glance returns a one-row data.frame with | | |
| nrow | number of rows | |
| ncol | number of columns | |
| complete.obs | number of rows that have no missing values | |
| na.fraction | fraction of values across all rows and columns that are missing | |

Author(s)

David Robinson, Benjamin Nutter

Source

Skew and Kurtosis functions are adapted from implementations in the moments package: Lukasz Komsta and Frederick Novomestky (2015). moments: Moments, cumulants, skewness, kurtosis and related tests. R package version 0.14. https://CRAN.R-project.org/package=moments

See Also

Other deprecated: bootstrap(), confint_tidy(), finish_glance(), fix_data_frame(), summary_tidiers, tidy.density(), tidy.dist(), tidy.ftable(), tidy.numeric()

Other deprecated: bootstrap(), confint_tidy(), finish_glance(), fix_data_frame(), summary_tidiers, tidy.density(), tidy.dist(), tidy.ftable(), tidy.numeric()

Other deprecated: bootstrap(), confint_tidy(), finish_glance(), fix_data_frame(), summary_tidiers, tidy.density(), tidy.dist(), tidy.ftable(), tidy.numeric()

```
td <- tidy(mtcars)
td
glance(mtcars)
library(ggplot2)
# compare mean and standard deviation
ggplot(td, aes(mean, sd)) + geom_point() +
    geom_text(aes(label = column), hjust = 1, vjust = 1) +</pre>
```

```
scale_x_log10() + scale_y_log10() + geom_abline()
```

durbinWatsonTest_tidiers

Tidy/glance a(n) durbinWatsonTest object

Description

For models that have only a single component, the tidy() and glance() methods are identical. Please see the documentation for both of those methods.

Usage

S3 method for class 'durbinWatsonTest'
tidy(x, ...)

S3 method for class 'durbinWatsonTest'
glance(x, ...)

Arguments

| х | An object of class durbinWatsonTest created by a call to car::durbinWatsonTest(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |
| | |

Value

A tibble::tibble() with columns:

| alternative | Alternative hypothesis (character). | |
|-----------------|---|--|
| autocorrelation | | |
| | Autocorrelation. | |
| p.value | The two-sided p-value associated with the observed statistic. | |
| statistic | Test statistic for Durbin-Watson test. | |
| method | Always 'Durbin-Watson Test'. | |

finish_glance

See Also

tidy(), glance(), car::durbinWatsonTest()
Other car tidiers: leveneTest_tidiers

Examples

```
# load modeling library
library(car)
# fit model
dw <- durbinWatsonTest(lm(mpg ~ wt, data = mtcars))
# summarize model fit with tidiers
tidy(dw)
# same output for all durbinWatsonTests
glance(dw)</pre>
```

| finish_glance | (Deprecated) Add logLik, AIC, BIC, and other common measurements |
|---------------|--|
| | to a glance of a prediction |

Description

This function is now deprecated in favor of using custom logic and the appropriate nobs() method.

Usage

finish_glance(ret, x)

Arguments

| ret | a one-row data frame (a partially complete glance) |
|-----|--|
| х | the prediction model |

Value

a one-row data frame with additional columns added, such as

| logLik | log likelihoods |
|-------------|--------------------------------|
| AIC | Akaike Information Criterion |
| BIC | Bayesian Information Criterion |
| deviance | deviance |
| df.residual | residual degrees of freedom |

See Also

```
Other deprecated: bootstrap(), confint_tidy(), data.frame_tidiers, fix_data_frame(), summary_tidiers, tidy.density(), tidy.dist(), tidy.ftable(), tidy.numeric()
```

fix_data_frame Ensure an object is a data frame, with rownames moved into a column

Description

This function is deprecated as of broom 0.7.0 and will be removed from a future release. Please see tibble::as_tibble.

Usage

fix_data_frame(x, newnames = NULL, newcol = "term")

Arguments

| Х | a data.frame or matrix |
|----------|--|
| newnames | new column names, not including the rownames |
| newcol | the name of the new rownames column |

Value

a data.frame, with rownames moved into a column and new column names assigned

See Also

Other deprecated: bootstrap(), confint_tidy(), data.frame_tidiers, finish_glance(), summary_tidiers, tidy.density(), tidy.dist(), tidy.ftable(), tidy.numeric()

glance.aareg Glance at a(n) aareg object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

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glance.aareg

Usage

```
## S3 method for class 'aareg'
glance(x, ...)
```

Arguments

| x | An aareg object returned from <pre>survival::aareg().</pre> |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| df | Degrees of freedom used by the model. |
|-----------|--|
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| statistic | Test statistic. |

See Also

glance(), survival::aareg()

Other aareg tidiers: tidy.aareg()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.cch(), glance.coxph(),
glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(), glance.survreg(),
tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(),
tidy.survfit(), tidy.survreg()
```

```
# load libraries for models and data
library(survival)
# fit model
afit <- aareg(
  Surv(time, status) ~ age + sex + ph.ecog,
 data = lung,
  dfbeta = TRUE
)
```

```
# summarize model fit with tidiers
tidy(afit)
```

glance.anova

Glance at a(n) anova object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'anova'
glance(x, ...)
```

Arguments

| x | An anova object, such as those created by stats::anova(), car::Anova(), car::leveneTest(), or car::linearHypothesis(). |
|-------|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |
| Value | |

A tibble::tibble() with exactly one row and columns:

| deviance | Deviance of the model. |
|-------------|------------------------------|
| df.residual | Residual degrees of freedom. |

glance.aov

Note

Note that the output of glance.anova() will vary depending on the initializing anova call. In some cases, it will just return an empty data frame. In other cases, glance.anova() may return columns that are also common to tidy.anova(). This is partly to preserve backwards compatibility with early versions of broom, but also because the underlying anova model yields components that could reasonably be interpreted as goodness-of-fit summaries too.

See Also

glance()

Other anova tidiers: glance.aov(), tidy.TukeyHSD(), tidy.anova(), tidy.aovlist(), tidy.aov(), tidy.manova()

Examples

```
# fit models
a <- lm(mpg ~ wt + qsec + disp, mtcars)
b <- lm(mpg ~ wt + qsec, mtcars)
mod <- anova(a, b)
# summarize model fit with tidiers
tidy(mod)
glance(mod)
# car::linearHypothesis() example
library(car)
mod_lht <- linearHypothesis(a, "wt - disp")
tidy(mod_lht)
glance(mod_lht)
```

glance.aov

Glance at a(n) lm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA. Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'aov'
glance(x, ...)
```

Arguments

```
x
...
```

An aov object, such as those created by stats::aov().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|----------|---|
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |

Note

Note that tidy.aov() now contains the numerator and denominator degrees of freedom, which were included in the output of glance.aov() in some previous versions of the package.

See Also

glance()

Other anova tidiers: glance.anova(), tidy.TukeyHSD(), tidy.anova(), tidy.aovlist(), tidy.aov(), tidy.manova()

Examples

```
a <- aov(mpg ~ wt + qsec + disp, mtcars)
tidy(a)</pre>
```

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Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'Arima'
glance(x, ...)
```

Arguments

| x | An object of class Arima created by stats::arima(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|--------|---|
| BIC | Bayesian Information Criterion for the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| sigma | Estimated standard error of the residuals. |

See Also

stats::arima()

Other Arima tidiers: tidy.Arima()

Examples

```
# fit model
fit <- arima(lh, order = c(1, 0, 0))
# summarize model fit with tidiers
tidy(fit)
glance(fit)</pre>
```

glance.betamfx

Glance at a(n) betamfx object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'betamfx'
glance(x, ...)
```

Arguments

х

A betamfx object.

. . .

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

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- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

This glance method wraps glance.betareg() for mfx::betamfx() objects.

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-----------------|--|
| BIC | Bayesian Information Criterion for the model. |
| df.null | Degrees of freedom used by the null model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| pseudo.r.square | d |
| | Like the R squared statistic, but for situations when the R squared statistic isn't defined. |

See Also

```
glance.betareg(), mfx::betamfx()
```

Other mfx tidiers: augment.betamfx(), augment.mfx(), glance.mfx(), tidy.betamfx(), tidy.mfx()

Examples

```
library(mfx)
```

```
# Simulate some data
set.seed(12345)
n <- 1000
x <- rnorm(n)
# Beta outcome
y <- rbeta(n, shape1 = plogis(1 + 0.5 * x), shape2 = (abs(0.2 * x)))
# Use Smithson and Verkuilen correction
y <- (y * (n - 1) + 0.5) / n
d <- data.frame(y, x)
mod_betamfx <- betamfx(y ~ x | x, data = d)
tidy(mod_betamfx, conf.int = TRUE)</pre>
```

```
# Compare with the naive model coefficients of the equivalent betareg call (not run)
# tidy(betamfx(y ~ x | x, data = d), conf.int = TRUE)
augment(mod_betamfx)
glance(mod_betamfx)
```

glance.betareg Glance at a(n) betareg object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'betareg'
glance(x, ...)
```

Arguments

| х | A betareg object produced by a call to betareg::betareg(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

glance.betareg

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. | |
|------------------|--|--|
| BIC | Bayesian Information Criterion for the model. | |
| df.null | Degrees of freedom used by the null model. | |
| df.residual | Residual degrees of freedom. | |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. | |
| nobs | Number of observations used. | |
| pseudo.r.squared | | |
| | Like the R squared statistic, but for situations when the R squared statistic isn't defined. | |

See Also

glance(), betareg::betareg()

Examples

```
# load libraries for models and data
library(betareg)
# load dats
data("GasolineYield", package = "betareg")
# fit model
mod <- betareg(yield ~ batch + temp, data = GasolineYield)
mod
# summarize model fit with tidiers
tidy(mod)
tidy(mod, conf.int = TRUE)
tidy(mod, conf.int = TRUE, conf.level = .99)
augment(mod)
glance(mod)
```

glance.biglm

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'biglm'
glance(x, ...)
```

Arguments

| х | A biglm object created by a call to biglm::biglm() or biglm::bigglm(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-------------|--|
| deviance | Deviance of the model. |
| df.residual | Residual degrees of freedom. |
| nobs | Number of observations used. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |

glance.binDesign

See Also

glance(), biglm::biglm(), biglm::bigglm()
Other biglm tidiers: tidy.biglm()

Examples

```
# load modeling library
library(biglm)
# fit model -- linear regression
bfit <- biglm(mpg ~ wt + disp, mtcars)</pre>
# summarize model fit with tidiers
tidy(bfit)
tidy(bfit, conf.int = TRUE)
tidy(bfit, conf.int = TRUE, conf.level = .9)
glance(bfit)
# fit model -- logistic regression
bgfit <- bigglm(am ~ mpg, mtcars, family = binomial())</pre>
# summarize model fit with tidiers
tidy(bgfit)
tidy(bgfit, exponentiate = TRUE)
tidy(bgfit, conf.int = TRUE)
tidy(bgfit, conf.int = TRUE, conf.level = .9)
tidy(bgfit, conf.int = TRUE, conf.level = .9, exponentiate = TRUE)
glance(bgfit)
```

glance.binDesign Glance at a(n) binDesign object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA. Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'binDesign'
glance(x, ...)
```

Arguments ×

. . .

| A binGroup::binDesign object. |
|--|
| Additional arguments. Not used. Needed to match generic signature only. Cau- |
| tionary note: Misspelled arguments will be absorbed in, where they will be |
| ignored. If the misspelled argument has a default value, the default value will be |
| used. For example, if you pass conf.lvel = 0.9, all computation will proceed |
| using conf.level = 0.95. Two exceptions here are: |
| tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| power | Power achieved by the analysis. |
|---------------|---|
| n | Sample size used to achieve this power. |
| power.reached | Whether the desired power was reached. |
| maxit | Number of iterations performed. |

See Also

glance(), binGroup::binDesign()

Other bingroup tidiers: tidy.binDesign(), tidy.binWidth()

Examples

```
# load libraries for models and data
library(binGroup)
des <- binDesign(
   nmax = 300, delta = 0.06,
   p.hyp = 0.1, power = .8
)
glance(des)
```

glance.cch

```
tidy(des)
library(ggplot2)
ggplot(tidy(des), aes(n, power)) +
  geom_line()
```

glance.cch

Glance at a(n) cch object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'cch'
glance(x, ...)
```

Arguments

| x | An cch object returned from survival::cch(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will |

Value

A tibble::tibble() with exactly one row and columns:

| iter | Iterations of algorithm/fitting procedure completed. |
|---------|--|
| p.value | P-value corresponding to the test statistic. |
| rscore | Robust log-rank statistic |
| score | Score. |
| n | number of predictions |
| nevent | number of events |

See Also

glance(), survival::cch()

Other cch tidiers: glance.survfit(), tidy.cch()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(), glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(), tidy.survfit(), tidy.survreg()
```

Examples

```
# load libraries for models and data
library(survival)
```

```
# examples come from cch documentation
subcoh <- nwtco$in.subcohort
selccoh <- with(nwtco, rel == 1 | subcoh == 1)
ccoh.data <- nwtco[selccoh, ]
ccoh.data$subcohort <- subcoh[selccoh]</pre>
```

```
# central-lab histology
ccoh.data$histol <- factor(ccoh.data$histol, labels = c("FH", "UH"))
# tumour stage
ccoh.data$stage <- factor(ccoh.data$stage, labels = c("I", "II", "III", "IV"))
ccoh.data$age <- ccoh.data$age / 12 # age in years
# fit model
fit.ccP <- cch(Surv(edrel, rel) ~ stage + histol + age,
    data = ccoh.data,
    subcoh = ~subcohort, id = ~seqno, cohort.size = 4028
)
# summarize model fit with tidiers + visualization
tidy(fit.ccP)
```

coefficient plot

glance.clm

```
library(ggplot2)
ggplot(tidy(fit.ccP), aes(x = estimate, y = term)) +
 geom_point() +
 geom_errorbarh(aes(xmin = conf.low, xmax = conf.high), height = 0) +
 geom_vline(xintercept = 0)
```

glance.clm

Glance at a(n) clm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'clm'
glance(x, ...)
```

Arguments

| x | A clm object returned from ordinal::clm(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-------------|---|
| BIC | Bayesian Information Criterion for the model. |
| df.residual | Residual degrees of freedom. |
| edf | The effective degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |

See Also

tidy, ordinal::clm()

Other ordinal tidiers: augment.clm(), augment.polr(), glance.clmm(), glance.polr(), glance.svyolr(), tidy.clmm(), tidy.clm(), tidy.polr(), tidy.svyolr()

Examples

```
# load libraries for models and data
library(ordinal)
# fit model
fit <- clm(rating ~ temp * contact, data = wine)
# summarize model fit with tidiers
tidy(fit)
tidy(fit, conf.int = TRUE, conf.level = 0.9)
tidy(fit, conf.int = TRUE, conf.type = "Wald", exponentiate = TRUE)
glance(fit)
augment(fit, type.predict = "prob")
augment(fit, type.predict = "class")
# ...and again with another model specification
fit2 <- clm(rating ~ temp, nominal = ~contact, data = wine)
tidy(fit2)
glance(fit2)
```

glance.clmm

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'clmm'
glance(x, ...)
```

Arguments

| x | A clmm object returned from ordinal::clmm(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

AICAkaike's Information Criterion for the model.BICBayesian Information Criterion for the model.edfThe effective degrees of freedom.logLikThe log-likelihood of the model. [stats::logLik()] may be a useful reference.nobsNumber of observations used.

See Also

```
tidy, ordinal::clmm()
```

```
Other ordinal tidiers: augment.clm(), augment.polr(), glance.clm(), glance.polr(), glance.svyolr(),
tidy.clmm(), tidy.clm(), tidy.polr(), tidy.svyolr()
```

Examples

```
# load libraries for models and data
library(ordinal)
# fit model
fit <- clmm(rating ~ temp + contact + (1 | judge), data = wine)
# summarize model fit with tidiers
tidy(fit)
tidy(fit, conf.int = TRUE, conf.level = 0.9)
tidy(fit, conf.int = TRUE, exponentiate = TRUE)
glance(fit)
# ...and again with another model specification
fit2 <- clmm(rating ~ temp + (1 | judge), nominal = ~contact, data = wine)
tidy(fit2)
glance(fit2)
```

glance.coeftest Glance at a(n) coeftest object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

glance.coeftest

Usage

```
## S3 method for class 'coeftest'
glance(x, ...)
```

Arguments ×

. .

| A coeftest object returned from <pre>lmtest::coeftest().</pre> |
|---|
| Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| adj.r.squared | Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account. |
|---------------|--|
| AIC | Akaike's Information Criterion for the model. |
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df | Degrees of freedom used by the model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| sigma | Estimated standard error of the residuals. |
| statistic | Test statistic. |

Note

Because of the way that lmtest::coeftest() retains information about the underlying model object, the returned columns for glance.coeftest() will vary depending on the arguments. Specifically, four columns are returned regardless: "Loglik", "AIC", "BIC", and "nobs". Users can obtain additional columns (e.g. "r.squared", "df") by invoking the "save = TRUE" argument as part of lmtest::coeftest(). See examples.

As an aside, goodness-of-fit measures such as R-squared are unaffected by the presence of heteroskedasticity. For further discussion see, e.g. chapter 8.1 of Wooldridge (2016).

References

Wooldridge, Jeffrey M. (2016) Introductory econometrics: A modern approach. (6th edition). Nelson Education.

See Also

glance(), lmtest::coeftest()

Examples

```
# load libraries for models and data
library(lmtest)
m <- lm(dist ~ speed, data = cars)</pre>
coeftest(m)
tidy(coeftest(m))
tidy(coeftest(m, conf.int = TRUE))
# a very common workflow is to combine lmtest::coeftest with alternate
# variance-covariance matrices via the sandwich package. The lmtest
# tidiers support this workflow too, enabling you to adjust the standard
# errors of your tidied models on the fly.
library(sandwich)
# "HC3" (default) robust SEs
tidy(coeftest(m, vcov = vcovHC))
# "HC2" robust SEs
tidy(coeftest(m, vcov = vcovHC, type = "HC2"))
# N-W HAC robust SEs
tidy(coeftest(m, vcov = NeweyWest))
# the columns of the returned tibble for glance.coeftest() will vary
# depending on whether the coeftest object retains the underlying model.
# Users can control this with the "save = TRUE" argument of coeftest().
glance(coeftest(m))
glance(coeftest(m, save = TRUE))
```

glance.coxph

glance.coxph

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

S3 method for class 'coxph'
glance(x, ...)

Arguments

| х | A coxph object returned from survival::coxph(). |
|---|--|
| | For tidy(), additional arguments passed to $summary(x, \ldots)$. Otherwise ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|--------|---|
| BIC | Bayesian Information Criterion for the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| n | The total number of observations. |
| nevent | Number of events. |
| nobs | Number of observations used. |
| | |

See survival::coxph.object for additional column descriptions.

See Also

glance(), survival::coxph()

Other coxph tidiers: augment.coxph(), tidy.coxph()

Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(), glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(), tidy.survfit(), tidy.survreg()

Examples

```
# load libraries for models and data
library(survival)
# fit model
cfit <- coxph(Surv(time, status) ~ age + sex, lung)</pre>
# summarize model fit with tidiers
tidy(cfit)
tidy(cfit, exponentiate = TRUE)
lp <- augment(cfit, lung)</pre>
risks <- augment(cfit, lung, type.predict = "risk")</pre>
expected <- augment(cfit, lung, type.predict = "expected")</pre>
glance(cfit)
# also works on clogit models
resp <- levels(logan$occupation)</pre>
n <- nrow(logan)</pre>
indx <- rep(1:n, length(resp))</pre>
logan2 <- data.frame(</pre>
  logan[indx, ],
  id = indx,
  tocc = factor(rep(resp, each = n))
)
logan2$case <- (logan2$occupation == logan2$tocc)</pre>
cl <- clogit(case ~ tocc + tocc:education + strata(id), logan2)</pre>
tidy(cl)
glance(cl)
library(ggplot2)
ggplot(lp, aes(age, .fitted, color = sex)) +
  geom_point()
ggplot(risks, aes(age, .fitted, color = sex)) +
  geom_point()
ggplot(expected, aes(time, .fitted, color = sex)) +
  geom_point()
```

glance.crr

Glance at a(n) crr object

glance.crr

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'crr'
glance(x, ...)
```

Arguments х

. . .

| A crr object returned from cmprsk::crr(). |
|---|
| Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| converged | Logical indicating if the model fitting procedure was succesful and converged. |
|-----------|--|
| df | Degrees of freedom used by the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| statistic | Test statistic. |

See Also

glance(), cmprsk::crr() Other cmprsk tidiers: tidy.crr()

Examples

```
library(cmprsk)
# time to loco-regional failure (lrf)
lrf_time <- rexp(100)
lrf_event <- sample(0:2, 100, replace = TRUE)
trt <- sample(0:1, 100, replace = TRUE)
strt <- sample(1:2, 100, replace = TRUE)
# fit model
x <- crr(lrf_time, lrf_event, cbind(trt, strt))
# summarize model fit with tidiers
tidy(x, conf.int = TRUE)
glance(x)</pre>
```

glance.cv.glmnet Glance at a(n) cv.glmnet object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'cv.glmnet'
glance(x, ...)
```

Arguments

```
Х
```

A cv.glmnet object returned from glmnet::cv.glmnet().

glance.cv.glmnet

. . .

| Additional arguments. Not used. Needed to match generic signature only. Cau- |
|--|
| tionary note: Misspelled arguments will be absorbed in , where they will be |
| ignored. If the misspelled argument has a default value, the default value will be |
| used. For example, if you pass conf.lvel = 0.9, all computation will proceed |
| using conf.level = 0.95. Two exceptions here are: |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| lambda.1se | The value of the penalization parameter lambda that results in the sparsest model while remaining within one standard error of the minimum loss. |
|------------|--|
| lambda.min | The value of the penalization parameter lambda that achieved minimum loss as estimated by cross validation. |
| nobs | Number of observations used. |

See Also

glance(), glmnet::cv.glmnet()
Other glmnet tidiers: glance.glmnet(), tidy.cv.glmnet(), tidy.glmnet()

Examples

```
# load libraries for models and data
library(glmnet)
set.seed(27)
nobs <- 100
nvar <- 50
real <- 5
x <- matrix(rnorm(nobs * nvar), nobs, nvar)
beta <- c(rnorm(real, 0, 1), rep(0, nvar - real))
y <- c(t(beta) %*% t(x)) + rnorm(nvar, sd = 3)
cvfit1 <- cv.glmnet(x, y)
tidy(cvfit1)
glance(cvfit1)
library(ggplot2)
tidied_cv <- tidy(cvfit1)</pre>
```

```
glance_cv <- glance(cvfit1)</pre>
# plot of MSE as a function of lambda
g <- ggplot(tidied_cv, aes(lambda, estimate)) +</pre>
 geom_line() +
 scale_x_log10()
g
# plot of MSE as a function of lambda with confidence ribbon
g <- g + geom_ribbon(aes(ymin = conf.low, ymax = conf.high), alpha = .25)
g
# plot of MSE as a function of lambda with confidence ribbon and choices
# of minimum lambda marked
g <- g +
 geom_vline(xintercept = glance_cv$lambda.min) +
 geom_vline(xintercept = glance_cv$lambda.1se, lty = 2)
g
# plot of number of zeros for each choice of lambda
ggplot(tidied_cv, aes(lambda, nzero)) +
 geom_line() +
 scale_x_log10()
# coefficient plot with min lambda shown
tidied <- tidy(cvfit1$glmnet.fit)</pre>
ggplot(tidied, aes(lambda, estimate, group = term)) +
 scale_x_log10() +
 geom_line() +
 geom_vline(xintercept = glance_cv$lambda.min) +
 geom_vline(xintercept = glance_cv$lambda.1se, lty = 2)
```

glance.drc

Glance at a(n) drc object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

glance.drc

Usage

```
## S3 method for class 'drc'
glance(x, ...)
```

Arguments

| х | A drc object produced by a call to drc::drm(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will |

upp rgi be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-------------|---|
| BIC | Bayesian Information Criterion for the model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| AICc | AIC corrected for small samples |

See Also

glance(), drc::drm()

Other drc tidiers: augment.drc(), tidy.drc()

Examples

```
# load libraries for models and data
library(drc)
# fit model
mod <- drm(dead / total ~ conc, type,</pre>
  weights = total, data = selenium, fct = LL.2(), type = "binomial"
)
# summarize model fit with tidiers
tidy(mod)
tidy(mod, conf.int = TRUE)
```

glance.ergm

glance(mod)

augment(mod, selenium)

glance.ergm

Glance at a(n) ergm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

S3 method for class 'ergm'
glance(x, deviance = FALSE, mcmc = FALSE, ...)

Arguments

| х | An ergm object returned from a call to ergm::ergm(). |
|----------|---|
| deviance | Logical indicating whether or not to report null and residual deviance for the model, as well as degrees of freedom. Defaults to FALSE. |
| мсмс | Logical indicating whether or not to report MCMC interval, burn-in and sample size used to estimate the model. Defaults to FALSE. |
| | Additional arguments to pass to ergm::summary(). Cautionary note: Mis- specified arguments may be silently ignored. |

Value

glance.ergm returns a one-row tibble with the columns

| independence | Whether the model assumed dyadic independence |
|--------------|---|
| iterations | The number of MCMLE iterations performed before convergence |
| logLik | If applicable, the log-likelihood associated with the model |
| AIC | The Akaike Information Criterion |

glance.factanal

| BIC | The Bayesian Information Criterion | |
|--|---|--|
| If deviance = TRUE, and if the model supports it, the tibble will also contain the columns | | |
| null.deviance | The null deviance of the model | |
| df.null | The degrees of freedom of the null deviance | |
| residual.deviance | | |
| | The residual deviance of the model | |
| df.residual | The degrees of freedom of the residual deviance | |

See Also

glance(), ergm::ergm(), ergm::summary.ergm()
Other ergm tidiers: tidy.ergm()

glance.factanal Glance at a(n) factanal object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

S3 method for class 'factanal'
glance(x, ...)

Arguments

A factanal object created by stats::factanal().

x ...

Additional arguments. Not used. Needed to match generic signature only. Cau-

tionary note: Misspelled arguments will be absorbed in ..., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| converged | Logical indicating if the model fitting procedure was succesful and converged. |
|----------------|--|
| df | Degrees of freedom used by the model. |
| method | Which method was used. |
| n | The total number of observations. |
| n.factors | The number of fitted factors. |
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| statistic | Test statistic. |
| total.variance | Total cumulative proportion of variance accounted for by all factors. |

See Also

glance(), stats::factanal()
Other factanal tidiers: augment.factanal(), tidy.factanal()

Examples

```
set.seed(123)
# generate data
library(dplyr)
library(purrr)
m1 <- tibble(</pre>
  v1 = c(1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 3, 3, 3, 3, 3, 4, 5, 6),
  v^2 = c(1, 2, 1, 1, 1, 1, 2, 1, 2, 1, 3, 4, 3, 3, 3, 4, 6, 5),
 v3 = c(3, 3, 3, 3, 3, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 5, 4, 6),
  v4 = c(3, 3, 4, 3, 3, 1, 1, 2, 1, 1, 1, 1, 2, 1, 1, 5, 6, 4),
  v5 = c(1, 1, 1, 1, 1, 3, 3, 3, 3, 3, 1, 1, 1, 1, 1, 6, 4, 5),
  v6 = c(1, 1, 1, 2, 1, 3, 3, 3, 4, 3, 1, 1, 1, 2, 1, 6, 5, 4)
)
# new data
m2 <- map_dfr(m1, rev)</pre>
# factor analysis objects
fit1 <- factanal(m1, factors = 3, scores = "Bartlett")</pre>
fit2 <- factanal(m1, factors = 3, scores = "regression")</pre>
```

glance.felm

```
# tidying the object
tidy(fit1)
tidy(fit2)
# augmented dataframe
augment(fit1)
augment(fit2)
# augmented dataframe (with new data)
augment(fit1, data = m2)
augment(fit2, data = m2)
```

glance.felm

Glance at a(n) *felm object*

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'felm'
glance(x, ...)
```

Arguments

| х | A felm object returned from lfe::felm(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will |

 augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| adj.r.squared | Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account. |
|---------------|--|
| df | Degrees of freedom used by the model. |
| df.residual | Residual degrees of freedom. |
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| sigma | Estimated standard error of the residuals. |
| statistic | Test statistic. |

Examples

```
# load libraries for models and data
library(lfe)
# use built-in `airquality` dataset
head(airquality)
# no FEs; same as lm()
est0 <- felm(Ozone ~ Temp + Wind + Solar.R, airquality)</pre>
# summarize model fit with tidiers
tidy(est0)
augment(est0)
# add month fixed effects
est1 <- felm(Ozone ~ Temp + Wind + Solar.R | Month, airquality)</pre>
# summarize model fit with tidiers
tidy(est1)
tidy(est1, fe = TRUE)
augment(est1)
glance(est1)
# the "se.type" argument can be used to switch out different standard errors
# types on the fly. In turn, this can be useful exploring the effect of
# different error structures on model inference.
tidy(est1, se.type = "iid")
tidy(est1, se.type = "robust")
# add clustered SEs (also by month)
est2 <- felm(Ozone ~ Temp + Wind + Solar.R | Month | 0 | Month, airquality)</pre>
```

```
# summarize model fit with tidiers
tidy(est2, conf.int = TRUE)
tidy(est2, conf.int = TRUE, se.type = "cluster")
tidy(est2, conf.int = TRUE, se.type = "robust")
tidy(est2, conf.int = TRUE, se.type = "iid")
```

glance.fitdistr Glance at a(n) fitdistr object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'fitdistr'
glance(x, ...)
```

Arguments

| x | A fitdistr object returned by MASS::fitdistr(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|--------|---|
| BIC | Bayesian Information Criterion for the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |

See Also

tidy(), MASS::fitdistr()
Other fitdistr tidiers: tidy.fitdistr()

Examples

```
# load libraries for models and data
library(MASS)
# generate data
set.seed(2015)
x <- rnorm(100, 5, 2)
# fit models
fit <- fitdistr(x, dnorm, list(mean = 3, sd = 1))
# summarize model fit with tidiers
tidy(fit)
glance(fit)</pre>
```

glance.fixest Glance at a(n) fixest object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

glance.fixest

Usage

```
## S3 method for class 'fixest'
glance(x, ...)
```

Arguments

| х | A fixest object returned from any of the fixest estimators |
|---|---|
| | Additional arguments passed to summary and confint. Important arguments are se and cluster. Other arguments are dof, exact_dof, forceCovariance, and keepBounded. See summary.fixest. |

Value

A tibble::tibble() with exactly one row and columns:

| adj.r.squared | Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account. |
|-------------------------|--|
| AIC | Akaike's Information Criterion for the model. |
| BIC | Bayesian Information Criterion for the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs pseudo.r.square | Number of observations used. |
| | Like the R squared statistic, but for situations when the R squared statistic isn't defined. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| sigma | Estimated standard error of the residuals. |
| within.r.square | ed |
| | R squared within fixed-effect groups. |

Note

All columns listed below will be returned, but some will be NA, depending on the type of model estimated. sigma, r.squared, adj.r.squared, and within.r.squared will be NA for any model other than feols. pseudo.r.squared will be NA for feols.

Examples

```
# load libraries for models and data
library(fixest)
gravity <-
   feols(
       log(Euros) ~ log(dist_km) | Origin + Destination + Product + Year, trade
   )</pre>
```

```
tidy(gravity)
glance(gravity)
augment(gravity, trade)
# to get robust or clustered SEs, users can either:
# 1) specify the arguments directly in the `tidy()` call
tidy(gravity, conf.int = TRUE, cluster = c("Product", "Year"))
tidy(gravity, conf.int = TRUE, se = "threeway")
# 2) or, feed tidy() a summary.fixest object that has already accepted
# these arguments
gravity_summ <- summary(gravity, cluster = c("Product", "Year"))
tidy(gravity_summ, conf.int = TRUE)
# approach (1) is preferred.</pre>
```

glance.Gam

```
Glance at a(n) Gam object
```

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'Gam'
glance(x, ...)
```

Arguments

```
Х
```

A Gam object returned from a call to gam::gam().

glance.gam

. . .

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

Glance at gam objects created by calls to mgcv::gam() with glance.gam().

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-------------|---|
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df | Degrees of freedom used by the model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |

See Also

glance(), gam::gam()
Other gam tidiers: tidy.Gam()

glance.gam

Glance at a(n) gam object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'gam'
glance(x, ...)
```

Arguments

| Х | A gam object returned from a call to mgcv::gam(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a powdete argument if it will |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| adj.r.squared | Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account. |
|---------------|--|
| AIC | Akaike's Information Criterion for the model. |
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df | Degrees of freedom used by the model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| npar | Number of parameters in the model. |

See Also

glance(), mgcv::gam() Other mgcv tidiers: tidy.gam()

Examples

```
# load libraries for models and data
library(mgcv)
# fit model
g <- gam(mpg ~ s(hp) + am + qsec, data = mtcars)
```

glance.garch

```
# summarize model fit with tidiers
tidy(g)
tidy(g, parametric = TRUE)
glance(g)
augment(g)
```

glance.garch Tidy a(n) garch object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'garch'
glance(x, test = c("box-ljung-test", "jarque-bera-test"), ...)
```

Arguments

| х | A garch object returned by tseries::garch(). |
|------|--|
| test | Character specification of which hypothesis test to use. The garch function reports 2 hypothesis tests: Jarque-Bera to residuals and Box-Ljung to squared residuals. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|--------|---|
| BIC | Bayesian Information Criterion for the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |

| method | Which method was used. |
|-----------|---|
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| statistic | Test statistic. |
| parameter | Parameter field in the htest, typically degrees of freedom. |

See Also

glance(), tseries::garch(),[]
Other garch tidiers: tidy.garch()

glance.geeglm Glance at a(n) geeglm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'geeglm'
glance(x, ...)
```

Arguments

x ... A geeglm object returned from a call to geepack::geeglm().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

glance.glm

Value

A tibble::tibble() with exactly one row and columns:

| alpha | Estimated correlation parameter for geepack::geeglm. | |
|------------------|--|--|
| df.residual | Residual degrees of freedom. | |
| gamma | Estimated scale parameter for geepack::geeglm. | |
| max.cluster.size | | |
| | Max number of elements in clusters. | |
| n.clusters | Number of clusters. | |

See Also

glance(), geepack::geeglm()

Examples

```
# load modeling library
library(geepack)
# load data
data(state)
ds <- data.frame(state.region, state.x77)
# fit model
geefit <- geeglm(Income ~ Frost + Murder,
    id = state.region,
    data = ds,
    corstr = "exchangeable"
)
# summarize model fit with tidiers
tidy(geefit)
tidy(geefit, conf.int = TRUE)
```

glance.glm

Glance at a(n) glm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'glm'
glance(x, ...)
```

Arguments

| х | A glm object returned from stats::glm(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|---------------|---|
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df.null | Degrees of freedom used by the null model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| null.deviance | Deviance of the null model. |

See Also

stats::glm()

Other Im tidiers: augment.glm(), augment.lm(), glance.lm(), glance.summary.lm(), glance.svyglm(), tidy.glm(), tidy.lm.beta(), tidy.lm(), tidy.mlm(), tidy.summary.lm()

glance.glmnet

Examples

```
g <- glm(am ~ mpg, mtcars, family = "binomial")
glance(g)</pre>
```

glance.glmnet

Glance at a(n) glmnet object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'glmnet'
glance(x, ...)
```

Arguments ×

| Х | A glmnet object returned from glmnet::glmnet(). |
|---|--|
| | Additional arguments. Not used. Needed to match gene |

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| nobs | Number of observations used. |
|---------|--|
| npasses | Total passes over the data across all lambda values. |
| nulldev | Null deviance. |

See Also

glance(), glmnet::glmnet()
Other glmnet tidiers: glance.cv.glmnet(), tidy.cv.glmnet(), tidy.glmnet()

Examples

```
# load libraries for models and data
library(glmnet)
set.seed(2014)
x <- matrix(rnorm(100 * 20), 100, 20)
y <- rnorm(100)</pre>
fit1 <- glmnet(x, y)</pre>
# summarize model fit with tidiers + visualization
tidy(fit1)
glance(fit1)
library(dplyr)
library(ggplot2)
tidied <- tidy(fit1) %>% filter(term != "(Intercept)")
ggplot(tidied, aes(step, estimate, group = term)) +
  geom_line()
ggplot(tidied, aes(lambda, estimate, group = term)) +
  geom_line() +
  scale_x_log10()
ggplot(tidied, aes(lambda, dev.ratio)) +
  geom_line()
# works for other types of regressions as well, such as logistic
g2 <- sample(1:2, 100, replace = TRUE)
fit2 <- glmnet(x, g2, family = "binomial")</pre>
tidy(fit2)
```

glance.glmRob Glance at a(n) glmRob object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

glance.glmRob

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'glmRob'
glance(x, ...)
```

Arguments

| x | A glmRob object returned from robust::glmRob(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with exactly one row and columns:

| deviance | Deviance of the model. |
|---------------|--|
| df.residual | Residual degrees of freedom. |
| nobs | Number of observations used. |
| null.deviance | Deviance of the null model. |
| sigma | Estimated standard error of the residuals. |

See Also

robust::glmRob()

Other robust tidiers: augment.lmRob(), glance.lmRob(), tidy.glmRob(), tidy.lmRob()

Examples

```
# load libraries for models and data
library(robust)
# fit model
gm <- glmRob(am ~ wt, data = mtcars, family = "binomial")</pre>
# summarize model fit with tidiers
tidy(gm)
glance(gm)
```

glance.gmm

Glance at a(n) gmm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'gmm'
glance(x, ...)
```

Arguments

х

. .

| A gmm object returned from gmm::gmm(). |
|--|
| Additional arguments. Not used. Needed to match generic signature only. Cau- |
| tionary note: Misspelled arguments will be absorbed in, where they will be |
| ignored. If the misspelled argument has a default value, the default value will be |
| used. For example, if you pass conf.lvel = 0.9, all computation will proceed |
| using conf.level = 0.95. Two exceptions here are: |
| • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

glance.gmm

Value

A tibble::tibble() with exactly one row and columns:

| Degrees of freedom used by the model. |
|--|
| Residual degrees of freedom. |
| Number of observations used. |
| P-value corresponding to the test statistic. |
| Test statistic. |
| |

See Also

glance(), gmm::gmm()

Other gmm tidiers: tidy.gmm()

Examples

```
# load libraries for models and data
library(gmm)
# examples come from the "gmm" package
# CAPM test with GMM
data(Finance)
r <- Finance[1:300, 1:10]</pre>
rm <- Finance[1:300, "rm"]</pre>
rf <- Finance[1:300, "rf"]</pre>
z <- as.matrix(r - rf)</pre>
t <- nrow(z)
zm <- rm - rf
h \leq matrix(zm, t, 1)
res <- gmm(z \sim zm, x = h)
# tidy result
tidy(res)
tidy(res, conf.int = TRUE)
tidy(res, conf.int = TRUE, conf.level = .99)
# coefficient plot
library(ggplot2)
library(dplyr)
tidy(res, conf.int = TRUE) %>%
  mutate(variable = reorder(term, estimate)) %>%
  ggplot(aes(estimate, variable)) +
  geom_point() +
  geom_errorbarh(aes(xmin = conf.low, xmax = conf.high)) +
  geom_vline(xintercept = 0, color = "red", lty = 2)
```

```
# from a function instead of a matrix
g <- function(theta, x) {</pre>
  e <- x[, 2:11] - theta[1] - (x[, 1] - theta[1]) %*% matrix(theta[2:11], 1, 10)</pre>
  gmat <- cbind(e, e * c(x[, 1]))
  return(gmat)
}
x <- as.matrix(cbind(rm, r))</pre>
res_black <- gmm(g, x = x, t0 = rep(0, 11))</pre>
tidy(res_black)
tidy(res_black, conf.int = TRUE)
# APT test with Fama-French factors and GMM
f1 <- zm
f2 <- Finance[1:300, "hml"] - rf</pre>
f3 <- Finance[1:300, "smb"] - rf
h <- cbind(f1, f2, f3)
res2 <- gmm(z ~ f1 + f2 + f3, x = h)
td2 <- tidy(res2, conf.int = TRUE)
td2
# coefficient plot
td2 %>%
  mutate(variable = reorder(term, estimate)) %>%
  ggplot(aes(estimate, variable)) +
  geom_point() +
  geom_errorbarh(aes(xmin = conf.low, xmax = conf.high)) +
  geom_vline(xintercept = 0, color = "red", lty = 2)
```

glance.ivreg Glance at a(n) ivreg object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

glance.ivreg

Usage

```
## S3 method for class 'ivreg'
glance(x, diagnostics = FALSE, ...)
```

Arguments

| х | An ivreg object created by a call to AER::ivreg(). |
|-------------|---|
| diagnostics | Logical indicating whether or not to return the Wu-Hausman and Sargan diag- nostic information. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

This tidier currently only supports ivreg-classed objects outputted by the AER package. The ivreg package also outputs objects of class ivreg, and will be supported in a later release.

Value

A tibble::tibble() with exactly one row and columns:

| adj.r.squared | Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account. |
|---------------|--|
| df | Degrees of freedom used by the model. |
| df.residual | Residual degrees of freedom. |
| nobs | Number of observations used. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| sigma | Estimated standard error of the residuals. |
| statistic | Wald test statistic. |
| p.value | P-value for the Wald test. |

Note

Beginning 0.7.0, glance.ivreg returns statistics for the Wu-Hausman test for endogeneity and the Sargan test of overidentifying restrictions. Sargan test values are returned as NA if the number of instruments is not greater than the number of endogenous regressors.

See Also

glance(), AER::ivreg()
Other ivreg tidiers: augment.ivreg(), tidy.ivreg()

Examples

```
# load libraries for models and data
library(AER)
# load data
data("CigarettesSW", package = "AER")
# fit model
ivr <- ivreg(</pre>
 log(packs) ~ income | population,
 data = CigarettesSW,
 subset = year == "1995"
)
# summarize model fit with tidiers
tidy(ivr)
tidy(ivr, conf.int = TRUE)
tidy(ivr, conf.int = TRUE, instruments = TRUE)
augment(ivr)
augment(ivr, data = CigarettesSW)
augment(ivr, newdata = CigarettesSW)
glance(ivr)
```

glance.kmeans Glance at a(n) kmeans object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

glance.kmeans

Usage

```
## S3 method for class 'kmeans'
glance(x, ...)
```

Arguments

| х | A kmeans object created by stats::kmeans(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| tidy() methods will warn when supplied an exponentiate will be ignored. | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | ignored. If the misspelled argument has a default value, the default value will used. For example, if you pass conf.lvel = 0.9, all computation will proce using conf.level = 0.95. Two exceptions here are: tidy() methods will warn when supplied an exponentiate argument is will be ignored. augment() methods will warn when supplied a newdata argument if it w |

Value

A tibble::tibble() with exactly one row and columns:

| betweenss | The total between-cluster sum of squares. |
|--------------|--|
| iter | Iterations of algorithm/fitting procedure completed. |
| tot.withinss | The total within-cluster sum of squares. |
| totss | The total sum of squares. |

See Also

glance(), stats::kmeans()

Other kmeans tidiers: augment.kmeans(), tidy.kmeans()

Examples

```
library(cluster)
library(modeldata)
library(dplyr)
data(hpc_data)
x <- hpc_data[, 2:5]
fit <- pam(x, k = 4)
tidy(fit)
glance(fit)
augment(fit, x)</pre>
```

glance.lavaan

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'lavaan'
glance(x, ...)
```

Arguments

| Х | A lavaan object, such as those returned from $lavaan::cfa()$, and $lavaan::sem()$. |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A one-row tibble::tibble with columns:

| chisq | Model chi squared | |
|-----------------|---|--|
| npar | Number of parameters in the model | |
| rmsea | Root mean square error of approximation | |
| rmsea.conf.high | | |
| | 95 percent upper bound on RMSEA | |

glance.lavaan

| srmr | Standardised root mean residual |
|----------------|---|
| agfi | Adjusted goodness of fit |
| cfi | Comparative fit index |
| tli | Tucker Lewis index |
| AIC | Akaike information criterion |
| BIC | Bayesian information criterion |
| ngroups | Number of groups in model |
| nobs | Number of observations included |
| norig | Number of observation in the original dataset |
| nexcluded | Number of excluded observations |
| converged | Logical - Did the model converge |
| estimator | Estimator used |
| missing_method | Method for eliminating missing data |

For further recommendations on reporting SEM and CFA models see Schreiber, J. B. (2017). Update to core reporting practices in structural equation modeling. Research in Social and Administrative Pharmacy, 13(3), 634-643. https://doi.org/10.1016/j.sapharm.2016.06.006

See Also

glance(), lavaan::cfa(), lavaan::sem(), lavaan::fitmeasures()

Other lavaan tidiers: tidy.lavaan()

Examples

```
library(lavaan)
# fit model
cfa.fit <- cfa(
    "F =~ x1 + x2 + x3 + x4 + x5",
    data = HolzingerSwineford1939, group = "school"
)
# summarize model fit with tidiers
glance(cfa.fit)</pre>
```

glance.lm

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'lm'
glance(x, ...)
```

Arguments

| x | An lm object created by stats::lm(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| adj.r.squared | Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account. |
|---------------|--|
| AIC | Akaike's Information Criterion for the model. |
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df.residual | Residual degrees of freedom. |

glance.lm

| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
|-----------|---|
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| sigma | Estimated standard error of the residuals. |
| statistic | Test statistic. |
| df | The degrees for freedom from the numerator of the overall F-statistic. This is new in broom 0.7.0. Previously, this reported the rank of the design matrix, which is one more than the numerator degrees of freedom of the overall F-statistic. |

See Also

glance(), glance.summary.lm()

Other lm tidiers: augment.glm(), augment.lm(), glance.glm(), glance.summary.lm(), glance.svyglm(), tidy.glm(), tidy.lm.beta(), tidy.lm(), tidy.mlm(), tidy.summary.lm()

Examples

```
library(ggplot2)
library(dplyr)
mod <- lm(mpg ~ wt + qsec, data = mtcars)</pre>
tidy(mod)
glance(mod)
# coefficient plot
d <- tidy(mod, conf.int = TRUE)</pre>
ggplot(d, aes(estimate, term, xmin = conf.low, xmax = conf.high, height = 0)) +
  geom_point() +
  geom_vline(xintercept = 0, lty = 4) +
  geom_errorbarh()
# aside: There are tidy() and glance() methods for lm.summary objects too.
# this can be useful when you want to conserve memory by converting large lm
# objects into their leaner summary.lm equivalents.
s <- summary(mod)</pre>
tidy(s, conf.int = TRUE)
glance(s)
augment(mod)
augment(mod, mtcars, interval = "confidence")
# predict on new data
```

```
newdata <- mtcars %>%
 head(6) %>%
 mutate(wt = wt + 1)
augment(mod, newdata = newdata)
# ggplot2 example where we also construct 95% prediction interval
# simpler bivariate model since we're plotting in 2D
mod2 <- lm(mpg ~ wt, data = mtcars)</pre>
au <- augment(mod2, newdata = newdata, interval = "prediction")</pre>
ggplot(au, aes(wt, mpg)) +
 geom_point() +
 geom_line(aes(y = .fitted)) +
 geom_ribbon(aes(ymin = .lower, ymax = .upper), col = NA, alpha = 0.3)
# predict on new data without outcome variable. Output does not include .resid
newdata <- newdata %>%
 select(-mpg)
augment(mod, newdata = newdata)
au <- augment(mod, data = mtcars)</pre>
ggplot(au, aes(.hat, .std.resid)) +
 geom_vline(size = 2, colour = "white", xintercept = 0) +
 geom_hline(size = 2, colour = "white", yintercept = 0) +
 geom_point() +
 geom_smooth(se = FALSE)
plot(mod, which = 6)
ggplot(au, aes(.hat, .cooksd)) +
 geom_vline(xintercept = 0, colour = NA) +
 geom_abline(slope = seq(0, 3, by = 0.5), colour = "white") +
 geom_smooth(se = FALSE) +
 geom_point()
# column-wise models
a <- matrix(rnorm(20), nrow = 10)
b <- a + rnorm(length(a))</pre>
result <- lm(b ~ a)</pre>
tidy(result)
```

glance.lmodel2 Glance at a(n) lmodel2 object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'lmodel2'
glance(x, ...)
```

Arguments ×

| х | A lmodel2 object returned by lmodel2::lmodel2(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cau- |
| | tionary note: Misspelled arguments will be absorbed in, where they will be |
| | ignored. If the misspelled argument has a default value, the default value will be |
| | used. For example, if you pass conf.lvel = 0.9, all computation will proceed |
| | using conf.level = 0.95. Two exceptions here are: |
| | |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| nobs | Number of observations used. |
|-----------|--|
| p.value | P-value corresponding to the test statistic. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| theta | Angle between OLS lines ' $lm(y \sim x)$ ' and ' $lm(x \sim y)$ ' |
| Н | H statistic for computing confidence interval of major axis slope |

See Also

glance(), lmodel2::lmodel2()
Other Imodel2 tidiers: tidy.lmodel2()

Examples

```
# load libraries for models and data
library(lmodel2)
data(mod2ex2)
Ex2.res <- lmodel2(Prey ~ Predators, data = mod2ex2, "relative", "relative", 99)
Ex2.res
# summarize model fit with tidiers + visualization
tidy(Ex2.res)
glance(Ex2.res)
# this allows coefficient plots with ggplot2
library(ggplot2)
ggplot(tidy(Ex2.res), aes(estimate, term, color = method)) +
geom_point() +
geom_errorbarh(aes(xmin = conf.low, xmax = conf.high)) +
geom_errorbarh(aes(xmin = conf.low, xmax = conf.high)) +
geom_errorbarh(aes(xmin = conf.low, xmax = conf.high))
```

glance.lmRob

Glance at a(n) lmRob object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'lmRob'
glance(x, ...)
```

glance.lmRob

Arguments

| х | A lmRob object returned from robust::lmRob(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| deviance | Deviance of the model. |
|-------------|--|
| df.residual | Residual degrees of freedom. |
| nobs | Number of observations used. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| sigma | Estimated standard error of the residuals. |

See Also

robust::lmRob()

Other robust tidiers: augment.lmRob(), glance.glmRob(), tidy.glmRob(), tidy.lmRob()

Examples

```
# load modeling library
library(robust)
# fit model
m <- lmRob(mpg ~ wt, data = mtcars)
# summarize model fit with tidiers
tidy(m)
augment(m)
glance(m)
```

glance.lmrob

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'lmrob'
glance(x, ...)
```

Arguments

| х | A lmrob object returned from robustbase::lmrob(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

For tidiers for robust models from the MASS package see tidy.rlm().

Value

A tibble::tibble() with exactly one row and columns:

| df.residual | Residual degrees of freedom. |
|-------------|--|
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| sigma | Estimated standard error of the residuals. |

glance.margins

See Also

robustbase::lmrob()

Other robustbase tidiers: augment.glmrob(), augment.lmrob(), tidy.glmrob(), tidy.lmrob()

Examples

```
if (requireNamespace("robustbase", quietly = TRUE)) {
 # load libraries for models and data
 library(robustbase)
 data(coleman)
 set.seed(0)
 m <- lmrob(Y ~ ., data = coleman)</pre>
 tidy(m)
 augment(m)
 glance(m)
 data(carrots)
 Rfit <- glmrob(cbind(success, total - success) ~ logdose + block,</pre>
    family = binomial, data = carrots, method = "Mqle",
    control = glmrobMqle.control(tcc = 1.2)
 )
 tidy(Rfit)
 augment(Rfit)
}
```

glance.margins Glance at a(n) margins object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'margins'
glance(x, ...)
```

Arguments

| x | A margins object returned from margins::margins(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will |

Value

A tibble::tibble() with exactly one row and columns:

be ignored.

| adj.r.squared | Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account. |
|---------------|--|
| df | Degrees of freedom used by the model. |
| df.residual | Residual degrees of freedom. |
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| sigma | Estimated standard error of the residuals. |
| statistic | Test statistic. |

glance.Mclust

Glance at a(n) Mclust object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

glance.Mclust

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'Mclust'
glance(x, ...)
```

Arguments

| х | An Mclust object return from mclust::Mclust(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| BIC | Bayesian Information Criterion for the model. |
|--------|--|
| df | Degrees of freedom used by the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| model | A string denoting the model type with optimal BIC |
| G | Number mixture components in optimal model |
| hypvol | If the other model contains a noise component, the value of the hypervolume parameter. Otherwise 'NA'. |

Examples

load library for models and data
library(mclust)

load data manipulation libraries
library(dplyr)

```
library(tibble)
library(purrr)
library(tidyr)
set.seed(27)
centers <- tibble(</pre>
 cluster = factor(1:3),
 # number points in each cluster
 num_points = c(100, 150, 50),
 # x1 coordinate of cluster center
 x1 = c(5, 0, -3),
 # x2 coordinate of cluster center
 x^2 = c(-1, 1, -2)
)
points <- centers %>%
 mutate(
   x1 = map2(num_points, x1, rnorm),
   x2 = map2(num_points, x2, rnorm)
 ) %>%
 select(-num_points, -cluster) %>%
 unnest(c(x1, x2))
# fit model
m <- Mclust(points)</pre>
# summarize model fit with tidiers
tidy(m)
augment(m, points)
glance(m)
```

```
glance.mfx
```

Glance at a(n) mfx object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

glance.mfx

Usage

```
## S3 method for class 'mfx'
glance(x, ...)
## S3 method for class 'logitmfx'
glance(x, ...)
## S3 method for class 'negbinmfx'
glance(x, ...)
## S3 method for class 'poissonmfx'
glance(x, ...)
## S3 method for class 'probitmfx'
glance(x, ...)
```

Arguments

. . .

x A logitmfx, negbinmfx, poissonmfx, or probitmfx object. (Note that betamfx objects receive their own set of tidiers.)

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

This generic glance method wraps glance.glm() for applicable objects from the mfx package.

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|---------------|---|
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df.null | Degrees of freedom used by the null model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| null.deviance | Deviance of the null model. |
| | |

See Also

```
glance.glm(),mfx::logitmfx(),mfx::negbinmfx(),mfx::poissonmfx(),mfx::probitmfx()
Other mfx tidiers: augment.betamfx(), augment.mfx(), glance.betamfx(), tidy.betamfx(),
tidy.mfx()
```

Examples

```
# load libraries for models and data
library(mfx)
# get the marginal effects from a logit regression
mod_logmfx <- logitmfx(am ~ cyl + hp + wt, atmean = TRUE, data = mtcars)</pre>
tidy(mod_logmfx, conf.int = TRUE)
# compare with the naive model coefficients of the same logit call
tidy(
 glm(am ~ cyl + hp + wt, family = binomial, data = mtcars),
 conf.int = TRUE
)
augment(mod_logmfx)
glance(mod_logmfx)
# another example, this time using probit regression
mod_probmfx <- probitmfx(am ~ cyl + hp + wt, atmean = TRUE, data = mtcars)</pre>
tidy(mod_probmfx, conf.int = TRUE)
augment(mod_probmfx)
glance(mod_probmfx)
```

glance.mjoint Glance at a(n) mjoint object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

glance.mjoint

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'mjoint'
glance(x, ...)
```

Arguments

```
x
...
```

An mjoint object returned from joineRML::mjoint().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|----------|---|
| BIC | Bayesian Information Criterion for the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| sigma2_j | The square root of the estimated residual variance for the j-th longitudinal pro- |
| | cess |

See Also

glance(), joineRML::mjoint()

Other mjoint tidiers: tidy.mjoint()

Examples

broom only skips running these examples because the example models take a
while to generatethey should run just fine, though!
Not run:

```
# load libraries for models and data
library(joineRML)
```

```
# fit a joint model with bivariate longitudinal outcomes
data(heart.valve)
hvd <- heart.valve[!is.na(heart.valve$log.grad) &</pre>
  !is.na(heart.valve$log.lvmi) &
  heart.valve$num <= 50, ]</pre>
fit <- mjoint(</pre>
  formLongFixed = list(
    "grad" = log.grad ~ time + sex + hs,
    "lvmi" = log.lvmi ~ time + sex
  ),
  formLongRandom = list(
    "grad" = ~ 1 | num,
    "lvmi" = ~ time | num
 ),
  formSurv = Surv(fuyrs, status) ~ age,
  data = hvd,
  inits = list("gamma" = c(0.11, 1.51, 0.80)),
  timeVar = "time"
)
# extract the survival fixed effects
tidy(fit)
# extract the longitudinal fixed effects
tidy(fit, component = "longitudinal")
# extract the survival fixed effects with confidence intervals
tidy(fit, ci = TRUE)
# extract the survival fixed effects with confidence intervals based
# on bootstrapped standard errors
bSE <- bootSE(fit, nboot = 5, safe.boot = TRUE)</pre>
tidy(fit, boot_se = bSE, ci = TRUE)
# augment original data with fitted longitudinal values and residuals
hvd2 <- augment(fit)</pre>
# extract model statistics
glance(fit)
## End(Not run)
```

glance.mlogit

glance.mlogit

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'mlogit'
glance(x, ...)
```

Arguments х

. .

| an object returned from mlogit::mlogit(). |
|--|
| Additional arguments. Not used. Needed to match generic signature only. Cau- |
| tionary note: Misspelled arguments will be absorbed in, where they will be |
| ignored. If the misspelled argument has a default value, the default value will be |
| used. For example, if you pass conf.lvel = 0.9, all computation will proceed |
| using conf.level = 0.95. Two exceptions here are: |
| |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|--------|--|
| BIC | Bayesian Information Criterion for the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| rho2 | McFadden's rho squared with respect to a market shares (constants-only) model. |
| rho20 | McFadden's rho squared with respect to an equal shares (no information) model. |
| | |

See Also

glance(), mlogit::mlogit()

Other mlogit tidiers: augment.mlogit(), tidy.mlogit()

Examples

```
# load libraries for models and data
library(mlogit)
data("Fishing", package = "mlogit")
Fish <- dfidx(Fishing, varying = 2:9, shape = "wide", choice = "mode")</pre>
# fit model
m <- mlogit(mode ~ price + catch | income, data = Fish)</pre>
# summarize model fit with tidiers
tidy(m)
augment(m)
glance(m)
```

glance.muhaz

Glance at a(n) muhaz object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'muhaz'
glance(x, ...)
```

Arguments

A muhaz object returned by muhaz::muhaz(). х

```
. . .
```

Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in ..., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| max.hazard | Maximal estimated hazard. |
|------------|---|
| max.time | The maximum observed event or censoring time. |
| min.hazard | Minimal estimated hazard. |
| min.time | The minimum observed event or censoring time. |
| nobs | Number of observations used. |

See Also

glance(), muhaz::muhaz()

Other muhaz tidiers: tidy.muhaz()

Examples

```
# load libraries for models and data
library(muhaz)
library(survival)
# fit model
x <- muhaz(ovarian$futime, ovarian$fustat)
# summarize model fit with tidiers
tidy(x)
glance(x)
```

glance.multinom Glance at a(n) multinom object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'multinom'
glance(x, ...)
```

Arguments

| х | A multinom object returned from nnet::multinom(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|----------|---|
| deviance | Deviance of the model. |
| edf | The effective degrees of freedom. |
| nobs | Number of observations used. |

See Also

glance(), nnet::multinom()

Other multinom tidiers: tidy.multinom()

Examples

```
# load libraries for models and data
library(nnet)
library(MASS)
```

example(birthwt)

glance.negbin

```
bwt.mu <- multinom(low ~ ., bwt)
tidy(bwt.mu)
glance(bwt.mu)
# or, for output from a multinomial logistic regression
fit.gear <- multinom(gear ~ mpg + factor(am), data = mtcars)
tidy(fit.gear)
glance(fit.gear)</pre>
```

glance.negbin

Glance at a(n) negbin object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'negbin'
glance(x, ...)
```

Arguments

| X | |
|---|--|
| | |

A negbin object returned by MASS::glm.nb().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|---------------|---|
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df.null | Degrees of freedom used by the null model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| null.deviance | Deviance of the null model. |

See Also

glance(), MASS::glm.nb()
Other glm.nb tidiers: tidy.negbin()

Examples

```
# load libraries for models and data
library(MASS)
# fit model
r <- glm.nb(Days ~ Sex / (Age + Eth * Lrn), data = quine)
# summarize model fit with tidiers
tidy(r)
glance(r)</pre>
```

```
glance.nlrq
```

Glance at a(n) nlrq object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

glance.nlrq

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'nlrq'
glance(x, ...)
```

Arguments

```
х
```

A nlrq object returned from quantreg::nlrq().

• • •

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-------------|---|
| BIC | Bayesian Information Criterion for the model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| tau | Quantile. |

See Also

glance(), quantreg::nlrq()

Other quantreg tidiers: augment.nlrq(), augment.rqs(), augment.rq(), glance.rq(), tidy.nlrq(), tidy.rqs(), tidy.rq()

Examples

```
# load modeling library
library(quantreg)
# build artificial data with multiplicative error
set.seed(1)
dat <- NULL</pre>
```

glance.nls

```
dat$x <- rep(1:25, 20)
dat$y <- SSlogis(dat$x, 10, 12, 2) * rnorm(500, 1, 0.1)</pre>
# fit the median using nlrq
mod <- nlrq(y ~ SSlogis(x, Asym, mid, scal),</pre>
  data = dat, tau = 0.5, trace = TRUE
)
# summarize model fit with tidiers
tidy(mod)
glance(mod)
augment(mod)
```

glance.nls

Glance at a(n) nls object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'nls'
glance(x, ...)
```

Arguments

| х | An nls object returned from stats::nls(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will |

 augment() methods will warn when supplied a newdata argument if it will be ignored.

glance.nls

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-------------|---|
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df.residual | Residual degrees of freedom. |
| finTol | The achieved convergence tolerance. |
| isConv | Whether the fit successfully converged. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| sigma | Estimated standard error of the residuals. |

See Also

tidy, stats::nls()

Other nls tidiers: augment.nls(), tidy.nls()

Examples

```
# fit model
n <- nls(mpg ~ k * e^wt, data = mtcars, start = list(k = 1, e = 2))
# summarize model fit with tidiers + visualization
tidy(n)
augment(n)
glance(n)
library(ggplot2)
ggplot(augment(n), aes(wt, mpg)) +
geom_point() +
geom_line(aes(y = .fitted))
newdata <- head(mtcars)
newdata$wt <- newdata$wt + 1
augment(n, newdata = newdata)
```

glance.orcutt

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'orcutt'
glance(x, ...)
```

Arguments

| х | An orcutt object returned from orcutt::cochrane.orcutt(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| adj.r.squared | Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account. |
|----------------|--|
| dw.original | Durbin-Watson statistic of original fit. |
| dw.transformed | Durbin-Watson statistic of transformed fit. |
| nobs | Number of observations used. |

glance.pam

| number.interact | tion |
|-----------------|--|
| | Number of interactions. |
| p.value.origina | al |
| | P-value of original Durbin-Watson statistic. |
| p.value.transfo | ormed |
| | P-value of autocorrelation after transformation. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| rho | Spearman's rho autocorrelation |

See Also

glance(), orcutt::cochrane.orcutt()
Other orcutt tidiers: tidy.orcutt()

Examples

```
# load libraries for models and data
library(orcutt)
# fit model and summarize results
reg <- lm(mpg ~ wt + qsec + disp, mtcars)
tidy(reg)
co <- cochrane.orcutt(reg)
tidy(co)
glance(co)
```

glance.pam

Glance at a(n) pam object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'pam'
glance(x, ...)
```

Arguments

| х | An pam object returned from cluster::pam() |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

```
avg.silhouette.width
```

The average silhouette width for the dataset.

See Also

```
glance(), cluster::pam()
```

Other pam tidiers: augment.pam(), tidy.pam()

Examples

```
# load libraries for models and data
library(dplyr)
library(ggplot2)
library(cluster)
library(modeldata)
data(hpc_data)
x <- hpc_data[, 2:5]
p <- pam(x, k = 4)
# summarize model fit with tidiers + visualization
tidy(p)
glance(p)
augment(p, x)
augment(p, x) %>%
ggplot(aes(compounds, input_fields)) +
```

glance.plm

```
geom_point(aes(color = .cluster)) +
geom_text(aes(label = cluster), data = tidy(p), size = 10)
```

glance.plm

Glance at a(n) plm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'plm'
glance(x, ...)
```

Arguments

| х | |
|---|--|
| | |
| | |

A plm objected returned by plm::plm().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

adj.r.squared Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account.

deviance Deviance of the model.

| df.residual | Residual degrees of freedom. |
|-------------|--|
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| statistic | F-statistic |

See Also

glance(), plm::plm()

Other plm tidiers: augment.plm(), tidy.plm()

Examples

```
# load libraries for models and data
library(plm)
# load data
data("Produc", package = "plm")
# fit model
zz <- plm(log(gsp) ~ log(pcap) + log(pc) + log(emp) + unemp,
data = Produc, index = c("state", "year")
)
# summarize model fit with tidiers
summary(zz)
tidy(zz, conf.int = TRUE)
tidy(zz, conf.int = TRUE, conf.level = 0.9)
augment(zz)
glance(zz)
```

glance.poLCA Glance at a(n) poLCA object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

glance.poLCA

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'poLCA'
glance(x, ...)
```

Arguments

| х | A poLCA object returned from poLCA::poLCA(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-------------|--|
| BIC | Bayesian Information Criterion for the model. |
| chi.squared | The Pearson Chi-Square goodness of fit statistic for multiway tables. |
| df | Degrees of freedom used by the model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. $[stats::logLik()]\ may be a useful reference.$ |
| nobs | Number of observations used. |
| g.squared | The likelihood ratio/deviance statistic |

See Also

glance(), poLCA::poLCA()

Other poLCA tidiers: augment.poLCA(), tidy.poLCA()

Examples

```
# load libraries for models and data
library(poLCA)
library(dplyr)
# generate data
data(values)
f <- cbind(A, B, C, D) ~ 1
# fit model
M1 <- poLCA(f, values, nclass = 2, verbose = FALSE)
M1
# summarize model fit with tidiers + visualization
tidy(M1)
augment(M1)
glance(M1)
library(ggplot2)
ggplot(tidy(M1), aes(factor(class), estimate, fill = factor(outcome))) +
  geom_bar(stat = "identity", width = 1) +
  facet_wrap(~variable)
# three-class model with a single covariate.
data(election)
f2a <- cbind(
  MORALG, CARESG, KNOWG, LEADG, DISHONG, INTELG,
  MORALB, CARESB, KNOWB, LEADB, DISHONB, INTELB
) ~ PARTY
nes2a <- poLCA(f2a, election, nclass = 3, nrep = 5, verbose = FALSE)</pre>
td <- tidy(nes2a)</pre>
td
ggplot(td, aes(outcome, estimate, color = factor(class), group = class)) +
  geom_line() +
  facet_wrap(~variable, nrow = 2) +
  theme(axis.text.x = element_text(angle = 90, hjust = 1))
au <- augment(nes2a)</pre>
au
count(au, .class)
```

glance.polr

```
# if the original data is provided, it leads to NAs in new columns
# for rows that weren't predicted
au2 <- augment(nes2a, data = election)
au2
dim(au2)
```

glance.polr

Glance at a(n) polr object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'polr'
glance(x, ...)
```

Arguments

| Х | | |
|---|---|---|
| | | |
| | • | • |

A polr object returned from MASS::polr().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-------------|---|
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df.residual | Residual degrees of freedom. |
| edf | The effective degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |

See Also

tidy, MASS::polr()

```
Other ordinal tidiers: augment.clm(), augment.polr(), glance.clm(), glance.clm(), glance.svyolr(),
tidy.clmm(), tidy.clm(), tidy.polr(), tidy.svyolr()
```

Examples

```
# load libraries for models and data
library(MASS)
# fit model
fit <- polr(Sat ~ Infl + Type + Cont, weights = Freq, data = housing)
# summarize model fit with tidiers
tidy(fit, exponentiate = TRUE, conf.int = TRUE)
glance(fit)
augment(fit, type.predict = "class")
fit2 <- polr(factor(gear) ~ am + mpg + qsec, data = mtcars)
tidy(fit, p.values = TRUE)
```

glance.pyears Glance at a(n) pyears object

glance.pyears

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'pyears'
glance(x, ...)
```

Arguments

| Х | A pyears object returned from survival::pyears(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cau- |
| | tionary note: Misspelled arguments will be absorbed in, where they will be |
| | ignored. If the misspelled argument has a default value, the default value will be |
| | used. For example, if you pass conf.lvel = 0.9, all computation will proceed |
| | using conf.level = 0.95. Two exceptions here are: |
| | |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| nobs | Number of observations used. |
|----------|--|
| total | total number of person-years tabulated |
| offtable | total number of person-years off table |

See Also

glance(), survival::pyears()

Other pyears tidiers: tidy.pyears()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(),
glance.coxph(),glance.survdiff(),glance.survexp(),glance.survfit(),glance.survreg(),
tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(),
tidy.survfit(), tidy.survreg()
```

Examples

```
# load libraries for models and data
library(survival)
# generate and format data
temp.yr <- tcut(mgus$dxyr, 55:92, labels = as.character(55:91))</pre>
temp.age <- tcut(mgus$age, 34:101, labels = as.character(34:100))</pre>
ptime <- ifelse(is.na(mgus$pctime), mgus$futime, mgus$pctime)</pre>
pstat <- ifelse(is.na(mgus$pctime), 0, 1)</pre>
pfit <- pyears(Surv(ptime / 365.25, pstat) ~ temp.yr + temp.age + sex, mgus,</pre>
  data.frame = TRUE
)
# summarize model fit with tidiers
tidy(pfit)
glance(pfit)
# if data.frame argument is not given, different information is present in
# output
pfit2 <- pyears(Surv(ptime / 365.25, pstat) ~ temp.yr + temp.age + sex, mgus)</pre>
tidy(pfit2)
glance(pfit2)
```

glance.ridgelm

```
Glance at a(n) ridgelm object
```

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'ridgelm'
glance(x, ...)
```

glance.ridgelm

Arguments

| х | A ridgelm object returned from MASS:::lm.ridge(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

This is similar to the output of select.ridgelm, but it is returned rather than printed.

Value

A tibble::tibble() with exactly one row and columns:

| kHKB | modified HKB estimate of the ridge constant |
|-----------|---|
| kLW | modified L-W estimate of the ridge constant |
| lambdaGCV | choice of lambda that minimizes GCV |

See Also

glance(), MASS::select.ridgelm(), MASS::lm.ridge()
Other ridgelm tidiers: tidy.ridgelm()

```
# load libraries for models and data
library(MASS)
names(longley)[1] <- "y"
# fit model and summarizd results
fit1 <- lm.ridge(y ~ ., longley)
tidy(fit1)
fit2 <- lm.ridge(y ~ ., longley, lambda = seq(0.001, .05, .001))
td2 <- tidy(fit2)
g2 <- glance(fit2)
# coefficient plot
library(ggplot2)
ggplot(td2, aes(lambda, estimate, color = term)) +
```

```
geom_line()
# GCV plot
ggplot(td2, aes(lambda, GCV)) +
 geom_line()
# add line for the GCV minimizing estimate
ggplot(td2, aes(lambda, GCV)) +
 geom_line() +
 geom_vline(xintercept = g2$lambdaGCV, col = "red", lty = 2)
```

glance.rlm

Glance at a(n) rlm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'rlm'
glance(x, ...)
```

Arguments х

. . .

| An rlm object returned by MASS::rlm(). |
|---|
| Additional arguments. Not used. Needed to match generic signature only. Cau - |
| tionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed |
| using conf.level = 0.95. Two exceptions here are: |
| • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

glance.rma

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-----------|--|
| BIC | Bayesian Information Criterion for the model. |
| converged | Logical indicating if the model fitting procedure was succesful and converged. |
| deviance | Deviance of the model. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| sigma | Estimated standard error of the residuals. |

See Also

glance(), MASS::rlm()

Other rlm tidiers: augment.rlm(), tidy.rlm()

Examples

```
# load libraries for models and data
library(MASS)
# fit model
r <- rlm(stack.loss ~ ., stackloss)
# summarize model fit with tidiers
tidy(r)
augment(r)
glance(r)
```

glance.rma

Glance at a(n) rma object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA. Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

S3 method for class 'rma'
glance(x, ...)

Arguments

| x | An rma object such as those created by metafor::rma(), metafor::rma.uni(), metafor::rma.glmm(), metafor::rma.mh(), metafor::rma.mv(), or metafor::rma.peto(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| cochran.qe | In meta-analysis, test statistic for the Cochran's Q_e test of residual heterogene- ity. | |
|--------------------|---|--|
| cochran.qm | In meta-analysis, test statistic for the Cochran's Q_m omnibus test of coefficients. | |
| df.residual | Residual degrees of freedom. | |
| h.squared | Value of the H-Squared statistic. | |
| i.squared | Value of the I-Squared statistic. | |
| measure | The measure used in the meta-analysis. | |
| method | Which method was used. | |
| nobs | Number of observations used. | |
| p.value.cochran.qe | | |
| | In meta-analysis, p-value for the Cochran's Q_e test of residual heterogeneity. | |
| p.value.cochran.qm | | |
| | In meta-analysis, p-value for the Cochran's Q_m omnibus test of coefficients. | |
| tau.squared | In meta-analysis, estimated amount of residual heterogeneity. | |
| tau.squared.se | In meta-analysis, standard error of residual heterogeneity. | |

glance.rq

Examples

```
library(metafor)

df <-
    escalc(
    measure = "RR",
    ai = tpos,
    bi = tneg,
    ci = cpos,
    di = cneg,
    data = dat.bcg
)

meta_analysis <- rma(yi, vi, data = df, method = "EB")
glance(meta_analysis)</pre>
```

glance.rq

Glance at a(n) rq object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'rq'
glance(x, ...)
```

Arguments

```
Х
```

An rq object returned from quantreg::rq().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

Only models with a single tau value may be passed. For multiple values, please use a purr::map() workflow instead, e.g.

```
taus %>%
map(function(tau_val) rq(y ~ x, tau = tau_val)) %>%
map_dfr(glance)
```

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-------------|---|
| BIC | Bayesian Information Criterion for the model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| tau | Quantile. |

See Also

glance(), quantreg::rq()

```
Other quantreg tidiers: augment.nlrq(), augment.rqs(), augment.rq(), glance.nlrq(), tidy.nlrq(),
tidy.rqs(), tidy.rq()
```

Examples

```
# load modeling library and data
library(quantreg)
data(stackloss)
# median (l1) regression fit for the stackloss data.
mod1 <- rq(stack.loss ~ stack.x, .5)
# weighted sample median
mod2 <- rq(rnorm(50) ~ 1, weights = runif(50))</pre>
```

. . .

glance.sarlm

```
# summarize model fit with tidiers
tidy(mod1)
glance(mod1)
augment(mod1)
tidy(mod2)
glance(mod2)
augment(mod2)
# varying tau to generate an rqs object
mod3 <- rq(stack.loss ~ stack.x, tau = c(.25, .5))
tidy(mod3)
augment(mod3)
# glance cannot handle rqs objects like `mod3`--use a purrr
# `map`-based workflow instead
```

glance.sarlm

Glance at a(*n*) *spatialreg object*

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'sarlm'
glance(x, ...)
```

Arguments

Х

An object returned from spatialreg::lagsarlm() or spatialreg::errorsarlm().

| | | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
|----|-----------------|---|
| | | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | | augment() methods will warn when supplied a newdata argument if it will be ignored. |
| Va | lue | |
| | A tibble::tibbl | e() with exactly one row and columns: |
| | AIC | Akaike's Information Criterion for the model. |
| | BIC | Bayesian Information Criterion for the model. |

- deviance Deviance of the model.
- logLik The log-likelihood of the model. [stats::logLik()] may be a useful reference.
- nobs Number of observations used.

See Also

glance(), spatialreg::lagsarlm(), spatialreg::errorsarlm(), spatialreg::sacsarlm()
Other spatialreg tidiers: augment.sarlm(), tidy.sarlm()

```
# load libraries for models and data
library(spatialreg)
library(spdep)
# load data
data(oldcol, package = "spdep")
listw <- nb2listw(COL.nb, style = "W")
# fit model
crime_sar <-
lagsarlm(CRIME ~ INC + HOVAL,
    data = COL.OLD,
    listw = listw,
    method = "eigen"
)
# summarize model fit with tidiers
tidy(crime_sar)
```

glance.smooth.spline

```
tidy(crime_sar, conf.int = TRUE)
glance(crime_sar)
augment(crime_sar)
# fit another model
crime_sem <- errorsarlm(CRIME ~ INC + HOVAL, data = COL.OLD, listw)</pre>
# summarize model fit with tidiers
tidy(crime_sem)
tidy(crime_sem, conf.int = TRUE)
glance(crime_sem)
augment(crime_sem)
# fit another model
crime_sac <- sacsarlm(CRIME ~ INC + HOVAL, data = COL.OLD, listw)</pre>
# summarize model fit with tidiers
tidy(crime_sac)
tidy(crime_sac, conf.int = TRUE)
glance(crime_sac)
augment(crime_sac)
```

glance.smooth.spline Tidy a(n) smooth.spine object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'smooth.spline'
glance(x, ...)
```

Arguments

| х | |
|---|--|
| | |

A smooth.spline object returned from stats::smooth.spline().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| crit | Minimized criterion |
|----------|---|
| cv.crit | Cross-validation score |
| df | Degrees of freedom used by the model. |
| lambda | Choice of lambda corresponding to 'spar'. |
| nobs | Number of observations used. |
| pen.crit | Penalized criterion. |
| spar | Smoothing parameter. |

See Also

augment(), stats::smooth.spline()

Other smoothing spline tidiers: augment.smooth.spline()

Examples

```
# fit model
spl <- smooth.spline(mtcars$wt, mtcars$mpg, df = 4)
# summarize model fit with tidiers
augment(spl, mtcars)
# calls original columns x and y
augment(spl)
library(ggplot2)
ggplot(augment(spl, mtcars), aes(wt, mpg)) +
geom_point() +
geom_line(aes(y = .fitted))</pre>
```

glance.speedglm Glance at a(n) speedglm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

glance.speedglm

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'speedglm'
glance(x, ...)
```

Arguments

| x | A speedglm object returned from speedglm::speedglm(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|---------------|---|
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df.null | Degrees of freedom used by the null model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| null.deviance | Deviance of the null model. |

See Also

speedglm::speedlm()

Other speedlm tidiers: augment.speedlm(), glance.speedlm(), tidy.speedglm(), tidy.speedlm()

Examples

```
# load libraries for models and data
library(speedglm)
# generate data
clotting <- data.frame(
    u = c(5, 10, 15, 20, 30, 40, 60, 80, 100),
    lot1 = c(118, 58, 42, 35, 27, 25, 21, 19, 18)
)
# fit model
fit <- speedglm(lot1 ~ log(u), data = clotting, family = Gamma(log))
# summarize model fit with tidiers
tidy(fit)
glance(fit)</pre>
```

glance.speedlm Glance at a(n) speedlm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'speedlm'
glance(x, ...)
```

Arguments

Х

A speedlm object returned from speedglm::speedlm().

glance.speedlm

. . .

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . ., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| adj.r.squared | Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account. |
|---------------|--|
| AIC | Akaike's Information Criterion for the model. |
| BIC | Bayesian Information Criterion for the model. |
| deviance | Deviance of the model. |
| df | Degrees of freedom used by the model. |
| df.residual | Residual degrees of freedom. |
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| statistic | F-statistic. |
| | |

See Also

speedglm::speedlm()

Other speedlm tidiers: augment.speedlm(), glance.speedglm(), tidy.speedglm(), tidy.speedlm()

```
# load modeling library
library(speedglm)
# fit model
mod <- speedlm(mpg ~ wt + qsec, data = mtcars, fitted = TRUE)
# summarize model fit with tidiers
tidy(mod)
glance(mod)
augment(mod)</pre>
```

glance.summary.lm Glance at a(n) summary.lm object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'summary.lm'
glance(x, ...)
```

Arguments

```
x
...
```

An lm object created by stats::lm().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

The glance.summary.lm() method is a potentially useful alternative to glance.lm(). For instance, if users have already converted large lm objects into their leaner summary.lm equivalents to conserve memory. Note, however, that this method does not return all of the columns of the non-summary method (e.g. AIC and BIC will be missing.)

Value

A tibble::tibble() with exactly one row and columns:

| adj.r.squared | Adjusted R squared statistic, which is like the R squared statistic except taking degrees of freedom into account. |
|---------------|---|
| df.residual | Residual degrees of freedom. |
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| r.squared | R squared statistic, or the percent of variation explained by the model. Also known as the coefficient of determination. |
| sigma | Estimated standard error of the residuals. |
| statistic | Test statistic. |
| df | The degrees for freedom from the numerator of the overall F-statistic. This is new in broom 0.7.0. Previously, this reported the rank of the design matrix, which is one more than the numerator degrees of freedom of the overall F-statistic. |

See Also

glance(), glance.summary.lm()

Other Im tidiers: augment.glm(), augment.lm(), glance.glm(), glance.lm(), glance.svyglm(), tidy.glm(), tidy.lm.beta(), tidy.lm(), tidy.mlm(), tidy.summary.lm()

```
library(ggplot2)
library(dplyr)
mod <- lm(mpg ~ wt + qsec, data = mtcars)</pre>
tidy(mod)
glance(mod)
# coefficient plot
d <- tidy(mod, conf.int = TRUE)</pre>
ggplot(d, aes(estimate, term, xmin = conf.low, xmax = conf.high, height = 0)) +
  geom_point() +
  geom_vline(xintercept = 0, lty = 4) +
  geom_errorbarh()
# aside: There are tidy() and glance() methods for lm.summary objects too.
# this can be useful when you want to conserve memory by converting large lm
# objects into their leaner summary.lm equivalents.
s <- summary(mod)</pre>
tidy(s, conf.int = TRUE)
```

```
glance(s)
augment(mod)
augment(mod, mtcars, interval = "confidence")
# predict on new data
newdata <- mtcars %>%
 head(6) %>%
 mutate(wt = wt + 1)
augment(mod, newdata = newdata)
# ggplot2 example where we also construct 95% prediction interval
# simpler bivariate model since we're plotting in 2D
mod2 <- lm(mpg ~ wt, data = mtcars)</pre>
au <- augment(mod2, newdata = newdata, interval = "prediction")</pre>
ggplot(au, aes(wt, mpg)) +
 geom_point() +
 geom_line(aes(y = .fitted)) +
 geom_ribbon(aes(ymin = .lower, ymax = .upper), col = NA, alpha = 0.3)
# predict on new data without outcome variable. Output does not include .resid
newdata <- newdata %>%
 select(-mpg)
augment(mod, newdata = newdata)
au <- augment(mod, data = mtcars)</pre>
ggplot(au, aes(.hat, .std.resid)) +
 geom_vline(size = 2, colour = "white", xintercept = 0) +
 geom_hline(size = 2, colour = "white", yintercept = 0) +
 geom_point() +
 geom_smooth(se = FALSE)
plot(mod, which = 6)
ggplot(au, aes(.hat, .cooksd)) +
 geom_vline(xintercept = 0, colour = NA) +
 geom_abline(slope = seq(0, 3, by = 0.5), colour = "white") +
 geom_smooth(se = FALSE) +
 geom_point()
# column-wise models
a <- matrix(rnorm(20), nrow = 10)
b <- a + rnorm(length(a))</pre>
result <- lm(b ~ a)</pre>
tidy(result)
```

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'survdiff'
glance(x, ...)
```

Arguments

Х

| An survdiff object returned from survival::survdiff(). | |
|---|--|
| Additional arguments. Not used. Needed to match generic signa | |

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| df | Degrees of freedom used by the model. |
|-----------|--|
| p.value | P-value corresponding to the test statistic. |
| statistic | Test statistic. |

See Also

```
glance(), survival::survdiff()
```

Other survdiff tidiers: tidy.survdiff()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(), glance.coxph(), glance.pyears(), glance.survexp(), glance.survfit(), glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(), tidy.survfit(), tidy.survreg()
```

Examples

```
# load libraries for models and data
library(survival)
# fit model
s <- survdiff(
  Surv(time, status) ~ pat.karno + strata(inst),
  data = lung
)
# summarize model fit with tidiers
tidy(s)
glance(s)</pre>
```

glance.survexp Glance at a(n) survexp object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'survexp'
glance(x, ...)
```

glance.survexp

Arguments

| х | An survexp object returned from survival::survexp(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| n.max | Maximum number of subjects at risk. |
|------------|-------------------------------------|
| n.start | Initial number of subjects at risk. |
| timepoints | Number of timepoints. |

See Also

glance(), survival::survexp()

Other survexp tidiers: tidy.survexp()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(),
glance.coxph(), glance.pyears(), glance.survdiff(), glance.survfit(), glance.survreg(),
tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(),
tidy.survfit(), tidy.survreg()
```

```
# load libraries for models and data
library(survival)
# fit model
sexpfit <- survexp(
  futime ~ 1,
  rmap = list(
    sex = "male",
    year = accept.dt,
    age = (accept.dt - birth.dt)
  ),
  method = "conditional",
  data = jasa
)
```

```
# summarize model fit with tidiers
tidy(sexpfit)
glance(sexpfit)
```

glance.survfit Glance at a(n) survfit object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

S3 method for class 'survfit'
glance(x, ...)

Arguments

| х | An survfit object returned from survival::survfit(). |
|---|--|
| | Additional arguments passed to summary.survfit(). Important arguments include rmean. |

Value

A tibble::tibble() with exactly one row and columns:

| events | Number of events. |
|-----------------|--|
| n.max | Maximum number of subjects at risk. |
| n.start | Initial number of subjects at risk. |
| nobs | Number of observations used. |
| records | Number of observations |
| rmean | Restricted mean (see [survival::print.survfit()]). |
| rmean.std.error | |
| | Postriated mean standard arror |

Restricted mean standard error.

| conf.low | lower end of confidence interval on median |
|-----------|--|
| conf.high | upper end of confidence interval on median |
| median | median survival |

See Also

glance(), survival::survfit()

Other cch tidiers: glance.cch(), tidy.cch()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(), glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(), tidy.survfit(), tidy.survreg()
```

```
# load libraries for models and data
library(survival)
# fit model
cfit <- coxph(Surv(time, status) ~ age + sex, lung)</pre>
sfit <- survfit(cfit)</pre>
# summarize model fit with tidiers + visualization
tidy(sfit)
glance(sfit)
library(ggplot2)
ggplot(tidy(sfit), aes(time, estimate)) +
  geom_line() +
  geom_ribbon(aes(ymin = conf.low, ymax = conf.high), alpha = .25)
# multi-state
fitCI <- survfit(Surv(stop, status * as.numeric(event), type = "mstate") ~ 1,</pre>
  data = mgus1, subset = (start == 0)
)
td_multi <- tidy(fitCI)</pre>
td_multi
ggplot(td_multi, aes(time, estimate, group = state)) +
  geom_line(aes(color = state)) +
  geom_ribbon(aes(ymin = conf.low, ymax = conf.high), alpha = .25)
```

glance.survreg

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'survreg'
glance(x, ...)
```

Arguments

| x | An survreg object returned from survival::survreg(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-------------|--|
| BIC | Bayesian Information Criterion for the model. |
| df | Degrees of freedom used by the model. |
| df.residual | Residual degrees of freedom. |
| iter | Iterations of algorithm/fitting procedure completed. |

glance.survreg

| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
|-----------|---|
| nobs | Number of observations used. |
| p.value | P-value corresponding to the test statistic. |
| statistic | Chi-squared statistic. |

See Also

glance(), survival::survreg()

Other survreg tidiers: augment.survreg(), tidy.survreg()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(), glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(), tidy.survfit(), tidy.survreg()
```

```
# load libraries for models and data
library(survival)
# fit model
sr <- survreg(</pre>
  Surv(futime, fustat) ~ ecog.ps + rx,
  ovarian,
  dist = "exponential"
)
# summarize model fit with tidiers + visualization
tidy(sr)
augment(sr, ovarian)
glance(sr)
# coefficient plot
td <- tidy(sr, conf.int = TRUE)</pre>
library(ggplot2)
ggplot(td, aes(estimate, term)) +
  geom_point() +
  geom_errorbarh(aes(xmin = conf.low, xmax = conf.high), height = 0) +
  geom_vline(xintercept = 0)
```

```
glance.svyglm
```

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'svyglm'
glance(x, maximal = x, ...)
```

Arguments

| х | A svyglm object returned from survey::svyglm(). |
|---------|---|
| maximal | A svyglm object corresponding to the maximal model against which to com- pute the BIC. See Lumley and Scott (2015) for details. Defaults to x, which is equivalent to not using a maximal model. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with exactly one row and columns:

| AIC | Akaike's Information Criterion for the model. |
|-----|---|
| BIC | Bayesian Information Criterion for the model. |

deviance Deviance of the model.

| df.null | Degrees of freedom used by the null model. |
|---------------|--|
| df.residual | Residual degrees of freedom. |
| null.deviance | Deviance of the null model. |

References

Lumley T, Scott A (2015). AIC and BIC for modelling with complex survey data. *Journal of Survey Statistics and Methodology*, 3(1).

See Also

```
survey::svyglm(), stats::glm(), survey::anova.svyglm
```

Other Im tidiers: augment.glm(), augment.lm(), glance.glm(), glance.lm(), glance.summary.lm(), tidy.glm(), tidy.lm.beta(), tidy.lm(), tidy.mlm(), tidy.summary.lm()

```
# load libraries for models and data
library(survey)
set.seed(123)
data(api)
# survey design
dstrat <-
  svydesign(
   id = ~1,
    strata = ~stype,
    weights = ~pw,
    data = apistrat,
    fpc = ~fpc
  )
# model
m <- svyglm(</pre>
  formula = sch.wide ~ ell + meals + mobility,
  design = dstrat,
  family = quasibinomial()
)
glance(m)
```

glance.svyolr

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'svyolr'
glance(x, ...)
```

Arguments

| х | A svyolr object returned from survey::svyolr(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| df.residual | Residual degrees of freedom. |
|-------------|-----------------------------------|
| edf | The effective degrees of freedom. |
| nobs | Number of observations used. |

glance.varest

See Also

tidy, survey::svyolr()

```
Other ordinal tidiers: augment.clm(), augment.polr(), glance.clmm(), glance.clm(), glance.polr(),
tidy.clmm(), tidy.clm(), tidy.polr(), tidy.svyolr()
```

Examples

```
library(broom)
library(survey)

data(api)
dclus1 <- svydesign(id = ~dnum, weights = ~pw, data = apiclus1, fpc = ~fpc)
dclus1 <- update(dclus1, mealcat = cut(meals, c(0, 25, 50, 75, 100)))
m <- svyolr(mealcat ~ avg.ed + mobility + stype, design = dclus1)
m
tidy(m, conf.int = TRUE)</pre>
```

glance.varest Glance at a(n) varest object

Description

Glance accepts a model object and returns a tibble::tibble() with exactly one row of model summaries. The summaries are typically goodness of fit measures, p-values for hypothesis tests on residuals, or model convergence information.

Glance never returns information from the original call to the modeling function. This includes the name of the modeling function or any arguments passed to the modeling function.

Glance does not calculate summary measures. Rather, it farms out these computations to appropriate methods and gathers the results together. Sometimes a goodness of fit measure will be undefined. In these cases the measure will be reported as NA.

Glance returns the same number of columns regardless of whether the model matrix is rank-deficient or not. If so, entries in columns that no longer have a well-defined value are filled in with an NA of the appropriate type.

Usage

```
## S3 method for class 'varest'
glance(x, ...)
```

Arguments

| x | A varest object produced by a call to vars::VAR(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with exactly one row and columns:

| lag.order | Lag order. |
|-----------|---|
| logLik | The log-likelihood of the model. [stats::logLik()] may be a useful reference. |
| n | The total number of observations. |
| nobs | Number of observations used. |

See Also

glance(), vars::VAR()

```
# load libraries for models and data
library(vars)
# load data
data("Canada", package = "vars")
# fit models
mod <- VAR(Canada, p = 1, type = "both")
# summarize model fit with tidiers
tidy(mod)
glance(mod)</pre>
```

Description

Broom tidies a number of lists that are effectively S3 objects without a class attribute. For example, stats::optim(), svd() and interp::interp() produce consistent output, but because they do not have a class attribute, they cannot be handled by S3 dispatch.

These functions look at the elements of a list and determine if there is an appropriate tidying method to apply to the list. Those tidiers are implemented as functions of the form tidy_<function> or glance_<function> and are not exported (but they are documented!).

If no appropriate tidying method is found, they throw an error.

Usage

glance_optim(x, ...)

Arguments

x ...

| A list returned from stats::optim(). |
|--|
| Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: |
| tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with exactly one row and columns:

| convergence | Convergence code. |
|----------------|--------------------------------------|
| function.count | Number of calls to 'fn'. |
| gradient.count | Number of calls to 'gr'. |
| value | Minimized or maximized output value. |

See Also

glance(), stats::optim()

Other list tidiers: list_tidiers, tidy_irlba(), tidy_optim(), tidy_svd(), tidy_xyz()

Examples

```
f <- function(x) (x[1] - 2)<sup>2</sup> + (x[2] - 3)<sup>2</sup> + (x[3] - 8)<sup>2</sup>
o <- optim(c(1, 1, 1), f)
```

leveneTest_tidiers Tidy/glance a(n) leveneTest object

Description

For models that have only a single component, the tidy() and glance() methods are identical. Please see the documentation for both of those methods.

Usage

```
## S3 method for class 'leveneTest'
tidy(x, ...)
```

Arguments

| х | An object of class anova created by a call to car::leveneTest(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| df | Degrees of freedom used by this term in the model. |
|-------------|---|
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| df.residual | Residual degrees of freedom. |

See Also

tidy(),glance(),car::leveneTest()
Other car tidiers: durbinWatsonTest_tidiers

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list_tidiers

Examples

```
# load libraries for models and data
library(car)
data(Moore)
lt <- with(Moore, leveneTest(conformity, fcategory))
tidy(lt)
glance(lt)
```

list_tidiers

Tidying methods for lists / returned values that are not S3 objects

Description

Broom tidies a number of lists that are effectively S3 objects without a class attribute. For example, stats::optim(), base::svd() and interp::interp() produce consistent output, but because they do not have a class attribute, they cannot be handled by S3 dispatch.

Usage

S3 method for class 'list'
tidy(x, ...)
S3 method for class 'list'

glance(x, ...)

Arguments

| х | A list, potentially representing an object that can be tidied. |
|---|--|
| | Additionally, arguments passed to the tidying function. |

Details

These functions look at the elements of a list and determine if there is an appropriate tidying method to apply to the list. Those tidiers are themselves are implemented as functions of the form tidy_<function> or glance_<function> and are not exported (but they are documented!).

If no appropriate tidying method is found, throws an error.

See Also

Other list tidiers: glance_optim(), tidy_irlba(), tidy_optim(), tidy_svd(), tidy_xyz()

null_tidiers

Description

tidy(NULL), glance(NULL) and augment(NULL) all return an empty tibble::tibble. This empty tibble can be treated a tibble with zero rows, making it convenient to combine with other tibbles using functions like purrr::map_df() on lists of potentially NULL objects.

Usage

```
## S3 method for class '`NULL`'
tidy(x, ...)
## S3 method for class '`NULL`'
glance(x, ...)
## S3 method for class '`NULL`'
augment(x, ...)
```

Arguments

| Х | The value NULL. |
|---|----------------------------------|
| | Additional arguments (not used). |

Value

An empty tibble::tibble.

See Also

tibble::tibble

```
sp_tidiers
```

Tidy a(n) SpatialPolygonsDataFrame object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Note that the sf package now defines tidy spatial objects and is the recommended approach to spatial data. sp tidiers are now deprecated in favor of sf::st_as_sf() and coercion methods found in other packages. See https://r-spatial.org/r/2023/05/15/evolution4.html for more on migration from retiring spatial packages.

summary_tidiers

Usage

```
## S3 method for class 'SpatialPolygonsDataFrame'
tidy(x, region = NULL, ...)
## S3 method for class 'SpatialPolygons'
tidy(x, ...)
## S3 method for class 'Polygons'
tidy(x, ...)
## S3 method for class 'Polygon'
tidy(x, ...)
## S3 method for class 'SpatialLinesDataFrame'
tidy(x, ...)
## S3 method for class 'Lines'
tidy(x, ...)
## S3 method for class 'Line'
tidy(x, ...)
```

Arguments

| X | A SpatialPolygonsDataFrame, SpatialPolygons, Polygons, Polygon, SpatialLinesDataFrame, Lines or Line object. |
|--------|--|
| region | name of variable used to split up regions |
| | not used by this method |

summary_tidiers (Deprecated) Tidy summaryDefault objects

Description

Tidiers for summaryDefault objects have been deprecated as of broom 0.7.0 in favor of skimr::skim().

Usage

```
## S3 method for class 'summaryDefault'
tidy(x, ...)
## S3 method for class 'summaryDefault'
glance(x, ...)
```

Arguments

| Х | A summaryDefault object, created by calling summary() on a vector. |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A one-row tibble::tibble with columns:

| minimum | Minimum value in original vector. |
|---------|---|
| q1 | First quartile of original vector. |
| median | Median of original vector. |
| mean | Mean of original vector. |
| q3 | Third quartile of original vector. |
| maximum | Maximum value in original vector. |
| na | Number of NA values in original vector. Column present only when original vector had at least one NA entry. |

See Also

Other deprecated: bootstrap(), confint_tidy(), data.frame_tidiers, finish_glance(), fix_data_frame(), tidy.density(), tidy.dist(), tidy.ftable(), tidy.numeric()

Other deprecated: bootstrap(), confint_tidy(), data.frame_tidiers, finish_glance(), fix_data_frame(), tidy.density(), tidy.dist(), tidy.ftable(), tidy.numeric()

```
v <- rnorm(1000)
s <- summary(v)
s
tidy(s)
glance(s)
v2 <- c(v,NA)
tidy(summary(v2))</pre>
```

tidy.aareg

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'aareg'
tidy(x, ...)

Arguments ×

. . .

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

robust.se is only present when x was created with dfbeta = TRUE.

Value

A tibble::tibble() with columns:

| estimate | The estimated value of the regression term. |
|-----------|---|
| p.value | The two-sided p-value associated with the observed statistic. |
| robust.se | robust version of standard error estimate. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| Z | z score. |

tidy.acf

See Also

tidy(), survival::aareg()

Other aareg tidiers: glance.aareg()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(),
glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(),
glance.survreg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(),
tidy.survfit(), tidy.survreg()
```

Examples

```
# load libraries for models and data
library(survival)
# fit model
afit <- aareg(
  Surv(time, status) ~ age + sex + ph.ecog,
  data = lung,
  dfbeta = TRUE
)
# summarize model fit with tidiers
tidy(afit)</pre>
```

tidy.acf

Tidy a(n) acf object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'acf'
tidy(x, ...)
```

Arguments

Х

An acf object created by stats::acf(), stats::pacf() or stats::ccf().

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tidy.anova

... Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in ..., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| acf | Autocorrelation. |
|-----|------------------|
| lag | Lag values. |

See Also

```
tidy(), stats::acf(), stats::pacf(), stats::ccf()
```

Other time series tidiers: tidy.spec(), tidy.ts(), tidy.zoo()

Examples

tidy(acf(lh, plot = FALSE))
tidy(ccf(mdeaths, fdeaths, plot = FALSE))
tidy(pacf(lh, plot = FALSE))

tidy.anova

Tidy a(n) anova object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'anova'
tidy(x, ...)
```

Arguments

| х | An anova object, such as those created by stats::anova(), car::Anova(), car::leveneTest(), or car::linearHypothesis(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

The term column of an ANOVA table can come with leading or trailing whitespace, which this tidying method trims.

For documentation on the tidier for car::leveneTest() output, see tidy.leveneTest()

Value

A tibble::tibble() with columns:

| df | Degrees of freedom used by this term in the model. |
|-----------|---|
| meansq | Mean sum of squares. Equal to total sum of squares divided by degrees of freedom. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| sumsq | Sum of squares explained by this term. |
| term | The name of the regression term. |

See Also

tidy(), stats::anova(), car::Anova(), car::leveneTest()

Other anova tidiers: glance.anova(), glance.aov(), tidy.TukeyHSD(), tidy.aovlist(), tidy.aov(), tidy.manova()

```
# fit models
a <- lm(mpg ~ wt + qsec + disp, mtcars)
b <- lm(mpg ~ wt + qsec, mtcars)
mod <- anova(a, b)</pre>
```

tidy.aov

```
# summarize model fit with tidiers
tidy(mod)
glance(mod)
# car::linearHypothesis() example
library(car)
mod_lht <- linearHypothesis(a, "wt - disp")
tidy(mod_lht)
glance(mod_lht)</pre>
```

tidy.aov

Tidy a(n) aov object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'aov'
tidy(x, intercept = FALSE, ...)

Arguments

| x | An aov object, such as those created by stats::aov(). |
|-----------|--|
| intercept | A logical indicating whether information on the intercept ought to be included. Passed to stats::summary.aov(). |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

The term column of an ANOVA table can come with leading or trailing whitespace, which this tidying method trims.

For documentation on the tidier for car::leveneTest() output, see tidy.leveneTest()

See Also

tidy(), stats::aov()

Other anova tidiers: glance.anova(), glance.aov(), tidy.TukeyHSD(), tidy.anova(), tidy.aovlist(), tidy.manova()

Examples

a <- aov(mpg ~ wt + qsec + disp, mtcars) tidy(a)

tidy.aovlist Tidy a(n) aovlist object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'aovlist' tidy(x, ...)

Arguments

An aovlist objects, such as those created by stats::aov().

Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in ..., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

The term column of an ANOVA table can come with leading or trailing whitespace, which this tidying method trims.

For documentation on the tidier for car::leveneTest() output, see tidy.leveneTest()

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tidy.Arima

Value

A tibble::tibble() with columns:

| df | Degrees of freedom used by this term in the model. |
|-----------|---|
| meansq | Mean sum of squares. Equal to total sum of squares divided by degrees of freedom. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| stratum | The error stratum. |
| sumsq | Sum of squares explained by this term. |
| term | The name of the regression term. |

See Also

tidy(), stats::aov()

Other anova tidiers: glance.anova(), glance.aov(), tidy.TukeyHSD(), tidy.anova(), tidy.aov(), tidy.manova()

Examples

a <- aov(mpg ~ wt + qsec + Error(disp / am), mtcars)
tidy(a)</pre>

tidy.Arima

Tidy a(n) Arima object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'Arima'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| х | An object of class Arima created by stats::arima(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

stats::arima()

Other Arima tidiers: glance.Arima()

```
# fit model
fit <- arima(lh, order = c(1, 0, 0))
# summarize model fit with tidiers
tidy(fit)
glance(fit)</pre>
```

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'betamfx'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| x | A betamfx object. |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will |

 augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

The mfx package provides methods for calculating marginal effects for various generalized linear models (GLMs). Unlike standard linear models, estimated model coefficients in a GLM cannot be directly interpreted as marginal effects (i.e., the change in the response variable predicted after a one unit change in one of the regressors). This is because the estimated coefficients are multiplicative, dependent on both the link function that was used for the estimation and any other variables that were included in the model. When calculating marginal effects, users must typically choose whether they want to use i) the average observation in the data, or ii) the average of the sample marginal effects. See vignette("mfxarticle") from the mfx package for more details.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- |
| | zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| atmean | TRUE if the marginal effects were originally calculated as the partial effects |
| | for the average observation. If FALSE, then these were instead calculated as |
| | average partial effects. |

See Also

```
tidy.betareg(), mfx::betamfx()
```

Other mfx tidiers: augment.betamfx(), augment.mfx(), glance.betamfx(), glance.mfx(), tidy.mfx()

Examples

```
library(mfx)
# Simulate some data
set.seed(12345)
n <- 1000
x <- rnorm(n)</pre>
# Beta outcome
y <- rbeta(n, shape1 = plogis(1 + 0.5 * x), shape2 = (abs(0.2 * x)))</pre>
# Use Smithson and Verkuilen correction
y <- (y * (n - 1) + 0.5) / n
d <- data.frame(y, x)</pre>
mod_betamfx <- betamfx(y ~ x | x, data = d)
tidy(mod_betamfx, conf.int = TRUE)
# Compare with the naive model coefficients of the equivalent betareg call (not run)
# tidy(betamfx(y ~ x | x, data = d), conf.int = TRUE)
augment(mod_betamfx)
glance(mod_betamfx)
```

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tidy.betareg

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'betareg'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| x | A betareg object produced by a call to betareg::betareg(). |
|------------|--|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

The tibble has one row for each term in the regression. The component column indicates whether a particular term was used to model either the "mean" or "precision". Here the precision is the inverse of the variance, often referred to as phi. At least one term will have been used to model the precision phi.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |

| estimate | The estimated value of the regression term. |
|-----------|---|
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| component | Whether a particular term was used to model the mean or the precision in the regression. See details. |

See Also

tidy(), betareg::betareg()

Examples

```
# load libraries for models and data
library(betareg)
# load dats
data("GasolineYield", package = "betareg")
# fit model
mod <- betareg(yield ~ batch + temp, data = GasolineYield)
mod
# summarize model fit with tidiers
tidy(mod)
tidy(mod, conf.int = TRUE)
tidy(mod, conf.int = TRUE, conf.level = .99)
augment(mod)
glance(mod)
```

tidy.biglm

Tidy a(n) biglm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

tidy.biglm

Usage

```
## S3 method for class 'biglm'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)
```

Arguments

| x | A biglm object created by a call to biglm::biglm() or biglm::bigglm(). |
|--------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| | |

See Also

tidy(), biglm::biglm(), biglm::bigglm()
Other biglm tidiers: glance.biglm()

Examples

```
# load modeling library
library(biglm)
# fit model -- linear regression
bfit <- biglm(mpg ~ wt + disp, mtcars)</pre>
# summarize model fit with tidiers
tidy(bfit)
tidy(bfit, conf.int = TRUE)
tidy(bfit, conf.int = TRUE, conf.level = .9)
glance(bfit)
# fit model -- logistic regression
bgfit <- bigglm(am ~ mpg, mtcars, family = binomial())</pre>
# summarize model fit with tidiers
tidy(bgfit)
tidy(bgfit, exponentiate = TRUE)
tidy(bgfit, conf.int = TRUE)
tidy(bgfit, conf.int = TRUE, conf.level = .9)
tidy(bgfit, conf.int = TRUE, conf.level = .9, exponentiate = TRUE)
glance(bgfit)
```

tidy.binDesign Tidy a(n) binDesign object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'binDesign'
tidy(x, ...)
```

Arguments

Х

A binGroup::binDesign() object.

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tidy.binWidth

. . .

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| n | Number of trials in given iteration. |
|-------|--------------------------------------|
| power | Power achieved for given value of n. |

See Also

tidy(), binGroup::binDesign()
Other bingroup tidiers: glance.binDesign(), tidy.binWidth()

Examples

```
library(binGroup)
des <- binDesign(
    nmax = 300, delta = 0.06,
    p.hyp = 0.1, power = .8
)
glance(des)
tidy(des)
# the ggplot2 equivalent of plot(des)
library(ggplot2)
ggplot(tidy(des), aes(n, power)) +
    geom_line()</pre>
```

tidy.binWidth Tidy a(n) binWidth object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'binWidth'
tidy(x, ...)
```

Arguments

| x | A binGroup::binWidth() object. |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will |

be ignored. i supplied a newdata argu war

Value

A tibble::tibble() with columns:

| alternative | Alternative hypothesis (character). |
|-------------|--|
| ci.width | Expected width of confidence interval. |
| р | True proportion. |
| n | Total sample size |

See Also

```
tidy(), binGroup::binWidth()
```

Other bingroup tidiers: glance.binDesign(), tidy.binDesign()

```
# load libraries
library(binGroup)
# fit model
bw <- binWidth(100, .1)</pre>
bw
# summarize model fit with tidiers
tidy(bw)
```

tidy.boot

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'boot'
tidy(
    x,
    conf.int = FALSE,
    conf.level = 0.95,
    conf.method = c("perc", "bca", "basic", "norm"),
    exponentiate = FALSE,
    ...
)
```

Arguments

| х | A boot::boot() object. |
|--------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| conf.method | Passed to the type argument of boot::boot.ci(). Defaults to "perc". The allowed types are "perc", "basic", "bca", and "norm". Does not support "stud" or "all". |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

If weights were provided to the boot function, an estimate column is included showing the weighted bootstrap estimate, and the standard error is of that estimate.

If there are no original statistics in the "boot" object, such as with a call to tsboot with orig.t = FALSE, the original and statistic columns are omitted, and only estimate and std.error columns shown.

Value

A tibble::tibble() with columns:

| bias | Bias of the statistic. |
|-----------|--|
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| statistic | Original value of the statistic. |

See Also

tidy(), boot::boot(), boot::tsboot(), boot::boot.ci(), rsample::bootstraps()

Examples

```
# load modeling library
library(boot)
clotting <- data.frame(</pre>
  u = c(5, 10, 15, 20, 30, 40, 60, 80, 100),
  lot1 = c(118, 58, 42, 35, 27, 25, 21, 19, 18),
  lot2 = c(69, 35, 26, 21, 18, 16, 13, 12, 12)
)
# fit models
g1 <- glm(lot2 ~ log(u), data = clotting, family = Gamma)
bootfun <- function(d, i) {</pre>
  coef(update(g1, data = d[i, ]))
}
bootres <- boot(clotting, bootfun, R = 999)</pre>
# summarize model fits with tidiers
tidy(g1, conf.int = TRUE)
tidy(bootres, conf.int = TRUE)
```

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tidy.btergm

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

This method tidies the coefficients of a bootstrapped temporal exponential random graph model estimated with the **xergm**. It simply returns the coefficients and their confidence intervals.

Usage

S3 method for class 'btergm'
tidy(x, conf.level = 0.95, exponentiate = FALSE, ...)

Arguments

| Х | A btergm::btergm() object. |
|--------------|--|
| conf.level | Confidence level for confidence intervals. Defaults to 0.95. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), btergm::btergm()

Examples

```
library(btergm)
library(network)
set.seed(5)
# create 10 random networks with 10 actors
networks <- list()</pre>
for (i in 1:10) {
  mat <- matrix(rbinom(100, 1, .25), nrow = 10, ncol = 10)</pre>
  diag(mat) <- 0</pre>
  nw <- network(mat)</pre>
  networks[[i]] <- nw</pre>
}
# create 10 matrices as covariates
covariates <- list()</pre>
for (i in 1:10) {
  mat <- matrix(rnorm(100), nrow = 10, ncol = 10)</pre>
  covariates[[i]] <- mat</pre>
}
# fit the model
mod <- btergm(networks ~ edges + istar(2) + edgecov(covariates), R = 100)</pre>
# summarize model fit with tidiers
tidy(mod)
```

tidy.cch

Tidy a(n) cch object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'cch'
tidy(x, conf.level = 0.95, ...)
```

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tidy.cch

Arguments

| An cch object returned from survival::cch(). |
|---|
| confidence level for CI |
| Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will |
| |

 augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), survival::cch()

Other cch tidiers: glance.cch(), glance.survfit()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(),
glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(),
glance.survreg(), tidy.aareg(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survexp(),
tidy.survfit(), tidy.survreg()
```

```
# load libraries for models and data
library(survival)
# examples come from cch documentation
subsch <= nwtcotin subschort</pre>
```

```
subcoh <- nwtco$in.subcohort
selccoh <- with(nwtco, rel == 1 | subcoh == 1)
ccoh.data <- nwtco[selccoh, ]</pre>
```

```
ccoh.data$subcohort <- subcoh[selccoh]</pre>
# central-lab histology
ccoh.data$histol <- factor(ccoh.data$histol, labels = c("FH", "UH"))</pre>
# tumour stage
ccoh.data$stage <- factor(ccoh.data$stage, labels = c("I", "II", "III", "IV"))</pre>
ccoh.data$age <- ccoh.data$age / 12 # age in years</pre>
# fit model
fit.ccP <- cch(Surv(edrel, rel) ~ stage + histol + age,</pre>
 data = ccoh.data,
 subcoh = ~subcohort, id = ~seqno, cohort.size = 4028
)
# summarize model fit with tidiers + visualization
tidy(fit.ccP)
# coefficient plot
library(ggplot2)
ggplot(tidy(fit.ccP), aes(x = estimate, y = term)) +
 geom_point() +
 geom_errorbarh(aes(xmin = conf.low, xmax = conf.high), height = 0) +
 geom_vline(xintercept = 0)
```

tidy.cld

Tidy a(n) cld object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'cld'
tidy(x, ...)
```

Arguments

| Х | A cld object created by calling multcomp::cld() on a glht, confint.glht() or summary.glht() object. |
|-----|---|
| ••• | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be |
| | ignored. If the misspelled argument has a default value, the default value will be |

used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

| A tibble::tibble() with columns: | |
|----------------------------------|--|
| contrast | Levels being compared. |
| letters | Compact letter display denoting all pair-wise comparisons. |

See Also

tidy(), multcomp::cld(), multcomp::summary.glht(), multcomp::confint.glht(), multcomp::glht()
Other multcomp tidiers: tidy.confint.glht(), tidy.glht(), tidy.summary.glht()

```
# load libraries for models and data
library(multcomp)
library(ggplot2)
amod <- aov(breaks ~ wool + tension, data = warpbreaks)</pre>
wht <- glht(amod, linfct = mcp(tension = "Tukey"))</pre>
tidy(wht)
ggplot(wht, aes(lhs, estimate)) +
  geom_point()
CI <- confint(wht)</pre>
tidy(CI)
ggplot(CI, aes(lhs, estimate, ymin = lwr, ymax = upr)) +
  geom_pointrange()
tidy(summary(wht))
ggplot(mapping = aes(lhs, estimate)) +
  geom_linerange(aes(ymin = lwr, ymax = upr), data = CI) +
  geom_point(aes(size = p), data = summary(wht)) +
  scale_size(trans = "reverse")
cld <- cld(wht)</pre>
tidy(cld)
```

tidy.clm

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'clm'
tidy(
    x,
    conf.int = FALSE,
    conf.level = 0.95,
    conf.type = c("profile", "Wald"),
    exponentiate = FALSE,
    ...
)
```

Arguments

| х | A clm object returned from ordinal::clm(). |
|--------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| conf.type | Whether to use "profile" or "Wald" confidence intervals, passed to the type argument of ordinal::confint.clm(). Defaults to "profile". |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

tidy.clm

Details

In broom 0.7.0 the coefficient_type column was renamed to coef.type, and the contents were changed as well.

Note that intercept type coefficients correspond to alpha parameters, location type coefficients correspond to beta parameters, and scale type coefficients correspond to zeta parameters.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy, ordinal::clm(), ordinal::confint.clm()

Other ordinal tidiers: augment.clm(), augment.polr(), glance.clmm(), glance.clm(), glance.polr(), glance.svyolr(), tidy.clmm(), tidy.polr(), tidy.svyolr()

```
# load libraries for models and data
library(ordinal)
# fit model
fit <- clm(rating ~ temp * contact, data = wine)
# summarize model fit with tidiers
tidy(fit)
tidy(fit, conf.int = TRUE, conf.level = 0.9)
tidy(fit, conf.int = TRUE, conf.type = "Wald", exponentiate = TRUE)
glance(fit)
augment(fit, type.predict = "prob")
augment(fit, type.predict = "class")
# ...and again with another model specification
fit2 <- clm(rating ~ temp, nominal = ~contact, data = wine)
tidy(fit2)
```

glance(fit2)

tidy.clmm

Tidy a(n) clmm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'clmm'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)

Arguments

| х | A clmm object returned from ordinal::clmm(). |
|--------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |

tidy.clmm

| estimate | The estimated value of the regression term. |
|-----------|---|
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

Note

In broom 0.7.0 the coefficient_type column was renamed to coef.type, and the contents were changed as well.

Note that intercept type coefficients correspond to alpha parameters, location type coefficients correspond to beta parameters, and scale type coefficients correspond to zeta parameters.

See Also

tidy, ordinal::clmm(), ordinal::confint.clm()

Other ordinal tidiers: augment.clm(), augment.polr(), glance.clmm(), glance.clm(), glance.polr(), glance.svyolr(), tidy.clm(), tidy.polr(), tidy.svyolr()

```
# load libraries for models and data
library(ordinal)
# fit model
fit <- clmm(rating ~ temp + contact + (1 | judge), data = wine)
# summarize model fit with tidiers
tidy(fit)
tidy(fit, conf.int = TRUE, conf.level = 0.9)
tidy(fit, conf.int = TRUE, exponentiate = TRUE)
glance(fit)
# ...and again with another model specification
fit2 <- clmm(rating ~ temp + (1 | judge), nominal = ~contact, data = wine)
tidy(fit2)
glance(fit2)
```

tidy.coeftest

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'coeftest'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| х | A coeftest object returned from <pre>lmtest::coeftest().</pre> |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

tidy.confint.glht

See Also

tidy(), lmtest::coeftest()

Examples

```
# load libraries for models and data
library(lmtest)
m <- lm(dist ~ speed, data = cars)</pre>
coeftest(m)
tidy(coeftest(m))
tidy(coeftest(m, conf.int = TRUE))
# a very common workflow is to combine lmtest::coeftest with alternate
# variance-covariance matrices via the sandwich package. The lmtest
# tidiers support this workflow too, enabling you to adjust the standard
# errors of your tidied models on the fly.
library(sandwich)
# "HC3" (default) robust SEs
tidy(coeftest(m, vcov = vcovHC))
# "HC2" robust SEs
tidy(coeftest(m, vcov = vcovHC, type = "HC2"))
# N-W HAC robust SEs
tidy(coeftest(m, vcov = NeweyWest))
# the columns of the returned tibble for glance.coeftest() will vary
# depending on whether the coeftest object retains the underlying model.
# Users can control this with the "save = TRUE" argument of coeftest().
glance(coeftest(m))
glance(coeftest(m, save = TRUE))
```

tidy.confint.glht Tidy a(n) confint.glht object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'confint.glht'
tidy(x, ...)
```

Arguments ×

. . .

| A confint.glht object created by calling multcomp::confint.glht() on a |
|--|
| glht object created with multcomp::glht(). |

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| contrast | Levels being compared. |
| estimate | The estimated value of the regression term. |

See Also

tidy(), multcomp::confint.glht(), multcomp::glht()
Other multcomp tidiers: tidy.cld(), tidy.glht(), tidy.summary.glht()

```
# load libraries for models and data
library(multcomp)
library(ggplot2)
amod <- aov(breaks ~ wool + tension, data = warpbreaks)
wht <- glht(amod, linfct = mcp(tension = "Tukey"))
tidy(wht)
ggplot(wht, aes(lhs, estimate)) +
geom_point()
```

tidy.confusionMatrix

```
CI <- confint(wht)
tidy(CI)
ggplot(CI, aes(lhs, estimate, ymin = lwr, ymax = upr)) +
geom_pointrange()
tidy(summary(wht))
ggplot(mapping = aes(lhs, estimate)) +
geom_linerange(aes(ymin = lwr, ymax = upr), data = CI) +
geom_point(aes(size = p), data = summary(wht)) +
scale_size(trans = "reverse")
cld <- cld(wht)
tidy(cld)</pre>
```

tidy.confusionMatrix Tidy a(n) confusionMatrix object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'confusionMatrix'
tidy(x, by_class = TRUE, ...)

Arguments

| х | An object of class confusionMatrix created by a call to caret::confusionMatrix(). |
|----------|--|
| by_class | Logical indicating whether or not to show performance measures broken down by class. Defaults to TRUE. When by_class = FALSE only returns a tibble with accuracy, kappa, and McNemar statistics. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| class | The class under consideration. |
|-----------|--|
| conf.high | Upper bound on the confidence interval for the estimate. |
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| term | The name of the regression term. |
| p.value | P-value for accuracy and kappa statistics. |

See Also

tidy(), caret::confusionMatrix()

Examples

```
# load libraries for models and data
library(caret)
set.seed(27)
# generate data
two_class_sample1 <- as.factor(sample(letters[1:2], 100, TRUE))</pre>
two_class_sample2 <- as.factor(sample(letters[1:2], 100, TRUE))</pre>
two_class_cm <- confusionMatrix(</pre>
  two_class_sample1,
  two_class_sample2
)
# summarize model fit with tidiers
tidy(two_class_cm)
tidy(two_class_cm, by_class = FALSE)
# multiclass example
six_class_sample1 <- as.factor(sample(letters[1:6], 100, TRUE))</pre>
six_class_sample2 <- as.factor(sample(letters[1:6], 100, TRUE))</pre>
six_class_cm <- confusionMatrix(</pre>
  six_class_sample1,
  six_class_sample2
)
# summarize model fit with tidiers
tidy(six_class_cm)
tidy(six_class_cm, by_class = FALSE)
```

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tidy.coxph

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'coxph'
tidy(x, exponentiate = FALSE, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| х | A coxph object returned from survival::coxph(). |
|--------------|--|
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | For tidy(), additional arguments passed to $summary(x,)$. Otherwise ignored. |

Value

A tibble::tibble() with columns:

| estimate | The estimated value of the regression term. |
|-----------|--|
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- |
| | zero. |
| std.error | The standard error of the regression term. |

See Also

tidy(), survival::coxph()

Other coxph tidiers: augment.coxph(), glance.coxph()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(),
glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(),
glance.survreg(), tidy.aareg(), tidy.cch(), tidy.pyears(), tidy.survdiff(), tidy.survexp(),
tidy.survfit(), tidy.survreg()
```

Examples

```
# load libraries for models and data
library(survival)
# fit model
cfit <- coxph(Surv(time, status) ~ age + sex, lung)</pre>
# summarize model fit with tidiers
tidy(cfit)
tidy(cfit, exponentiate = TRUE)
lp <- augment(cfit, lung)</pre>
risks <- augment(cfit, lung, type.predict = "risk")</pre>
expected <- augment(cfit, lung, type.predict = "expected")</pre>
glance(cfit)
# also works on clogit models
resp <- levels(logan$occupation)</pre>
n <- nrow(logan)</pre>
indx <- rep(1:n, length(resp))</pre>
logan2 <- data.frame(</pre>
  logan[indx, ],
  id = indx,
  tocc = factor(rep(resp, each = n))
)
logan2$case <- (logan2$occupation == logan2$tocc)</pre>
cl <- clogit(case ~ tocc + tocc:education + strata(id), logan2)</pre>
tidy(cl)
glance(cl)
library(ggplot2)
ggplot(lp, aes(age, .fitted, color = sex)) +
  geom_point()
ggplot(risks, aes(age, .fitted, color = sex)) +
  geom_point()
ggplot(expected, aes(time, .fitted, color = sex)) +
  geom_point()
```

tidy.crr

Tidy a(n) cmprsk object

tidy.crr

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'crr'
tidy(x, exponentiate = FALSE, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| x | A crr object returned from cmprsk::crr(). |
|--------------|---|
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |

See Also

tidy(), cmprsk::crr()

Other cmprsk tidiers: glance.crr()

Examples

```
library(cmprsk)
```

```
# time to loco-regional failure (lrf)
lrf_time <- rexp(100)
lrf_event <- sample(0:2, 100, replace = TRUE)
trt <- sample(0:1, 100, replace = TRUE)
strt <- sample(1:2, 100, replace = TRUE)
# fit model
x <- crr(lrf_time, lrf_event, cbind(trt, strt))
# summarize model fit with tidiers
tidy(x, conf.int = TRUE)
glance(x)</pre>
```

tidy.cv.glmnet Tidy a(n) cv.glmnet object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'cv.glmnet'
tidy(x, ...)
```

Arguments

x A cv.glmnet object returned from glmnet::cv.glmnet().

... Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in ..., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

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- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| lambda | Value of penalty parameter lambda. |
|-----------|---|
| nzero | Number of non-zero coefficients for the given lambda. |
| std.error | The standard error of the regression term. |
| conf.low | lower bound on confidence interval for cross-validation estimated loss. |
| conf.high | upper bound on confidence interval for cross-validation estimated loss. |
| estimate | Median loss across all cross-validation folds for a given lamdba |

See Also

```
tidy(), glmnet::cv.glmnet()
```

Other glmnet tidiers: glance.cv.glmnet(), glance.glmnet(), tidy.glmnet()

```
# load libraries for models and data
library(glmnet)
set.seed(27)
nobs <- 100
nvar <- 50
real <- 5
x <- matrix(rnorm(nobs * nvar), nobs, nvar)</pre>
beta <- c(rnorm(real, 0, 1), rep(0, nvar - real))</pre>
y \leq c(t(beta) \% t(x)) + rnorm(nvar, sd = 3)
cvfit1 <- cv.glmnet(x, y)</pre>
tidy(cvfit1)
glance(cvfit1)
library(ggplot2)
tidied_cv <- tidy(cvfit1)</pre>
glance_cv <- glance(cvfit1)</pre>
# plot of MSE as a function of lambda
g <- ggplot(tidied_cv, aes(lambda, estimate)) +</pre>
```

```
geom_line() +
 scale_x_log10()
g
# plot of MSE as a function of lambda with confidence ribbon
g <- g + geom_ribbon(aes(ymin = conf.low, ymax = conf.high), alpha = .25)
g
# plot of MSE as a function of lambda with confidence ribbon and choices
# of minimum lambda marked
g <- g +
 geom_vline(xintercept = glance_cv$lambda.min) +
 geom_vline(xintercept = glance_cv$lambda.1se, lty = 2)
g
# plot of number of zeros for each choice of lambda
ggplot(tidied_cv, aes(lambda, nzero)) +
 geom_line() +
 scale_x_log10()
# coefficient plot with min lambda shown
tidied <- tidy(cvfit1$glmnet.fit)</pre>
ggplot(tidied, aes(lambda, estimate, group = term)) +
 scale_x_log10() +
 geom_line() +
 geom_vline(xintercept = glance_cv$lambda.min) +
 geom_vline(xintercept = glance_cv$lambda.1se, lty = 2)
```

tidy.density (Deprecated) Tidy density objects

Description

(Deprecated) Tidy density objects

Usage

```
## S3 method for class 'density'
tidy(x, ...)
```

Arguments

x A density object returned from stats::density().

... Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in ..., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

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tidy.dist

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble with two columns: points x where the density is estimated, and estimated density y.

See Also

Other deprecated: bootstrap(), confint_tidy(), data.frame_tidiers, finish_glance(), fix_data_frame(), summary_tidiers, tidy.dist(), tidy.ftable(), tidy.numeric()

tidy.dist (Deprecated) Tidy dist objects

Description

(Deprecated) Tidy dist objects

Usage

```
## S3 method for class 'dist'
tidy(x, diagonal = attr(x, "Diag"), upper = attr(x, "Upper"), ...)
```

Arguments

| х | A dist object returned from stats::dist(). |
|----------|---|
| diagonal | Logical indicating whether or not to tidy the diagonal elements of the distance matrix. Defaults to whatever was based to the diag argument of stats::dist(). |
| upper | Logical indicating whether or not to tidy the upper half of the distance matrix. Defaults to whatever was based to the upper argument of stats::dist(). |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Details

If the distance matrix does not include an upper triangle and/or diagonal, the tidied version will not either.

Value

A tibble::tibble with one row for each pair of items in the distance matrix, with columns:

| item1 | First item |
|----------|------------------------|
| item2 | Second item |
| distance | Distance between items |

See Also

```
Other deprecated: bootstrap(), confint_tidy(), data.frame_tidiers, finish_glance(), fix_data_frame(),
summary_tidiers, tidy.density(), tidy.ftable(), tidy.numeric()
```

Examples

```
cars_dist <- dist(t(mtcars[, 1:4]))
cars_dist
tidy(cars_dist)
tidy(cars_dist, upper = TRUE)
tidy(cars_dist, diagonal = TRUE)</pre>
```

tidy.drc

Tidy a(n) drc object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'drc'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| х | A drc object produced by a call to drc::drm(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |

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. . .

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

The tibble has one row for each curve and term in the regression. The curveid column indicates the curve.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| curve | Index identifying the curve. |

See Also

tidy(), drc::drm()

Other drc tidiers: augment.drc(), glance.drc()

```
# load libraries for models and data
library(drc)
# fit model
mod <- drm(dead / total ~ conc, type,
   weights = total, data = selenium, fct = LL.2(), type = "binomial"
)
# summarize model fit with tidiers
tidy(mod)</pre>
```

```
tidy(mod, conf.int = TRUE)
glance(mod)
augment(mod, selenium)
```

tidy.emmGrid Tid

Tidy a(n) emmGrid object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'emmGrid'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| х | An emmGrid object. |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments passed to emmeans::summary.emmGrid() or lsmeans::summary.ref.grid(). Cautionary note : misspecified arguments may be silently ignored! |

Details

Returns a data frame with one observation for each estimated marginal mean, and one column for each combination of factors. When the input is a contrast, each row will contain one estimated contrast.

There are a large number of arguments that can be passed on to emmeans::summary.emmGrid() or lsmeans::summary.ref.grid().

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |

```
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```

tidy.emmGrid

| df | Degrees of freedom used by this term in the model. |
|-----------|---|
| p.value | The two-sided p-value associated with the observed statistic. |
| std.error | The standard error of the regression term. |
| estimate | Expected marginal mean |
| statistic | T-ratio statistic |

See Also

```
tidy(), emmeans::ref_grid(), emmeans::emmeans(), emmeans::contrast()
Other emmeans tidiers: tidy.lsmobj(), tidy.ref.grid(), tidy.summary_emm()
```

```
# load libraries for models and data
library(emmeans)
# linear model for sales of oranges per day
oranges_lm1 <- lm(sales1 ~ price1 + price2 + day + store, data = oranges)</pre>
# reference grid; see vignette("basics", package = "emmeans")
oranges_rg1 <- ref_grid(oranges_lm1)</pre>
td <- tidy(oranges_rg1)</pre>
td
# marginal averages
marginal <- emmeans(oranges_rg1, "day")</pre>
tidy(marginal)
# contrasts
tidy(contrast(marginal))
tidy(contrast(marginal, method = "pairwise"))
# plot confidence intervals
library(ggplot2)
ggplot(tidy(marginal, conf.int = TRUE), aes(day, estimate)) +
  geom_point() +
  geom_errorbar(aes(ymin = conf.low, ymax = conf.high))
# by multiple prices
by_price <- emmeans(oranges_lm1, "day",</pre>
  by = "price2",
  at = list(
    price1 = 50, price2 = c(40, 60, 80),
    day = c("2", "3", "4")
 )
)
by_price
```

```
tidy(by_price)
ggplot(tidy(by_price, conf.int = TRUE), aes(price2, estimate, color = day)) +
  geom_line() +
  geom_errorbar(aes(ymin = conf.low, ymax = conf.high))
# joint_tests
tidy(joint_tests(oranges_lm1))
```

tidy.epi.2by2 *Tidy a(n) epi.2by2 object*

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'epi.2by2'
tidy(x, parameters = c("moa", "stat"), ...)

Arguments

| x | A epi.2by2 object produced by a call to epiR::epi.2by2() |
|------------|---|
| parameters | Return measures of association (moa) or test statistics (stat), default is moa (measures of association) |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

The tibble has a column for each of the measures of association or tests contained in massoc or massoc.detail when epiR::epi.2by2() is called.

tidy.ergm

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| df | Degrees of freedom used by this term in the model. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| term | The name of the regression term. |
| estimate | Estimated measure of association |

See Also

tidy(), epiR::epi.2by2()

```
# load libraries for models and data
library(epiR)
# generate data
dat <- matrix(c(13, 2163, 5, 3349), nrow = 2, byrow = TRUE)
rownames(dat) <- c("DF+", "DF-")
colnames(dat) <- c("FUS+", "FUS-")
# fit model
fit <- epi.2by2(
    dat = as.table(dat), method = "cross.sectional",
    conf.level = 0.95, units = 100, outcome = "as.columns"
)
# summarize model fit with tidiers
tidy(fit, parameters = "moa")
tidy(fit, parameters = "stat")
```

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

The methods should work with any model that conforms to the **ergm** class, such as those produced from weighted networks by the **ergm.count** package.

Usage

```
## S3 method for class 'ergm'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)
```

Arguments

| x | An ergm object returned from a call to ergm::ergm(). |
|--------------|--|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments to pass to ergm::summary(). Cautionary note: Misspecified arguments may be silently ignored. |

Value

A tibble::tibble with one row for each coefficient in the exponential random graph model, with columns:

| term | The term in the model being estimated and tested |
|------------|--|
| estimate | The estimated coefficient |
| std.error | The standard error |
| mcmc.error | The MCMC error |
| p.value | The two-sided p-value |

References

Hunter DR, Handcock MS, Butts CT, Goodreau SM, Morris M (2008b). **ergm**: A Package to Fit, Simulate and Diagnose Exponential-Family Models for Networks. *Journal of Statistical Software*, 24(3). https://www.jstatsoft.org/v24/i03/.

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tidy.factanal

See Also

tidy(), ergm::ergm(), ergm::control.ergm(), ergm::summary()

Other ergm tidiers: glance.ergm()

Examples

```
# load libraries for models and data
library(ergm)
# load the Florentine marriage network data
data(florentine)
# fit a model where the propensity to form ties between
# families depends on the absolute difference in wealth
gest <- ergm(flomarriage ~ edges + absdiff("wealth"))</pre>
# show terms, coefficient estimates and errors
tidy(gest)
# show coefficients as odds ratios with a 99% CI
tidy(gest, exponentiate = TRUE, conf.int = TRUE, conf.level = 0.99)
# take a look at likelihood measures and other
# control parameters used during MCMC estimation
glance(gest)
glance(gest, deviance = TRUE)
glance(gest, mcmc = TRUE)
```

tidy.factanal Tidy a(n) factanal object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'factanal'
tidy(x, ...)
```

Arguments

| A factanal object created by stats::factanal(). |
|---|
| Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| augment() methods will warn when supplied a newdata argument if it will be ignored. |
| |

Value

A tibble::tibble() with columns:

| variable | Variable under consideration. |
|------------|---|
| uniqueness | Proportion of residual, or unexplained variance |
| flX | Factor loading for level X. |

See Also

tidy(), stats::factanal()

Other factanal tidiers: augment.factanal(), glance.factanal()

```
set.seed(123)
# generate data
library(dplyr)
library(purrr)
m1 <- tibble(</pre>
  v1 = c(1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 3, 3, 3, 3, 3, 4, 5, 6),
 v^2 = c(1, 2, 1, 1, 1, 1, 2, 1, 2, 1, 3, 4, 3, 3, 3, 4, 6, 5),
 v3 = c(3, 3, 3, 3, 3, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 5, 4, 6),
 v4 = c(3, 3, 4, 3, 3, 1, 1, 2, 1, 1, 1, 1, 2, 1, 1, 5, 6, 4),
 v5 = c(1, 1, 1, 1, 1, 3, 3, 3, 3, 3, 1, 1, 1, 1, 1, 6, 4, 5),
  v6 = c(1, 1, 1, 2, 1, 3, 3, 3, 4, 3, 1, 1, 1, 2, 1, 6, 5, 4)
)
# new data
m2 <- map_dfr(m1, rev)</pre>
# factor analysis objects
fit1 <- factanal(m1, factors = 3, scores = "Bartlett")</pre>
fit2 <- factanal(m1, factors = 3, scores = "regression")</pre>
```

tidy.felm

```
# tidying the object
tidy(fit1)
tidy(fit2)
# augmented dataframe
augment(fit1)
augment(fit2)
# augmented dataframe (with new data)
augment(fit1, data = m2)
augment(fit2, data = m2)
```

tidy.felm

Tidy a(n) felm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'felm'
tidy(
    x,
    conf.int = FALSE,
    conf.level = 0.95,
    fe = FALSE,
    se.type = c("default", "iid", "robust", "cluster"),
    ...
)
```

Arguments

| x | A felm object returned from lfe::felm(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| fe | Logical indicating whether or not to include estimates of fixed effects. Defaults to FALSE. |

| se.type | Character indicating the type of standard errors. Defaults to using those of the underlying felm() model object, e.g. clustered errors for models that were provided a cluster specification. Users can override these defaults by specifying an appropriate alternative: "iid" (for homoskedastic errors), "robust" (for Eicker-Huber-White robust errors), or "cluster" (for clustered standard errors; if the model object supports it). |
|---------|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(),lfe::felm()

Other felm tidiers: augment.felm()

```
# load libraries for models and data
library(lfe)
# use built-in `airquality` dataset
head(airquality)
# no FEs; same as lm()
est0 <- felm(Ozone ~ Temp + Wind + Solar.R, airquality)</pre>
```

tidy.fitdistr

```
# summarize model fit with tidiers
tidy(est0)
augment(est0)
# add month fixed effects
est1 <- felm(Ozone ~ Temp + Wind + Solar.R | Month, airquality)</pre>
# summarize model fit with tidiers
tidy(est1)
tidy(est1, fe = TRUE)
augment(est1)
glance(est1)
# the "se.type" argument can be used to switch out different standard errors
# types on the fly. In turn, this can be useful exploring the effect of
# different error structures on model inference.
tidy(est1, se.type = "iid")
tidy(est1, se.type = "robust")
# add clustered SEs (also by month)
est2 <- felm(Ozone ~ Temp + Wind + Solar.R | Month | 0 | Month, airquality)
# summarize model fit with tidiers
tidy(est2, conf.int = TRUE)
tidy(est2, conf.int = TRUE, se.type = "cluster")
tidy(est2, conf.int = TRUE, se.type = "robust")
tidy(est2, conf.int = TRUE, se.type = "iid")
```

tidy.fitdistr Tidy a(n) fitdistr object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'fitdistr'
tidy(x, ...)
```

Arguments

```
Х
```

A fitdistr object returned by MASS::fitdistr().

| Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed |
|--|
| using conf.level = 0.95. Two exceptions here are:tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| • sugment() methods will warn when supplied a newdata argument if it will |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| estimate | The estimated value of the regression term. |
|-----------|---|
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), MASS::fitdistr()

Other fitdistr tidiers: glance.fitdistr()

```
# load libraries for models and data
library(MASS)
# generate data
set.seed(2015)
x <- rnorm(100, 5, 2)
# fit models
fit <- fitdistr(x, dnorm, list(mean = 3, sd = 1))
# summarize model fit with tidiers
tidy(fit)
glance(fit)</pre>
```

tidy.fixest

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'fixest'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| x | A fixest object returned from any of the fixest estimators |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments passed to summary and confint. Important arguments are seand cluster. Other arguments are dof, exact_dof, forceCovariance, and keepBounded. See summary.fixest. |

Details

The fixest package provides a family of functions for estimating models with arbitrary numbers of fixed-effects, in both an OLS and a GLM context. The package also supports robust (i.e. White) and clustered standard error reporting via the generic summary.fixest() command. In a similar vein, the tidy() method for these models allows users to specify a desired standard error correction either 1) implicitly via the supplied fixest object, or 2) explicitly as part of the tidy call. See examples below.

Note that fixest confidence intervals are calculated assuming a normal distribution – this assumes infinite degrees of freedom for the CI. (This assumption is distinct from the degrees of freedom used to calculate the standard errors. For more on degrees of freedom with clusters and fixed effects, see https://github.com/lrberge/fixest/issues/6 and https://github.com/sgaure/lfe/issues/1#issuecomment-530646990)

Value

A tibble::tibble() with columns:

conf.high Upper bound on the confidence interval for the estimate.

| conf.low | Lower bound on the confidence interval for the estimate. |
|-----------|---|
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(),fixest::feglm(),fixest::fenegbin(),fixest::feNmlm(),fixest::femlm(),fixest::feols(), fixest::fepois()

Other fixest tidiers: augment.fixest()

```
# load libraries for models and data
library(fixest)
gravity <-
  feols(
   log(Euros) ~ log(dist_km) | Origin + Destination + Product + Year, trade
  )
tidy(gravity)
glance(gravity)
augment(gravity, trade)
# to get robust or clustered SEs, users can either:
# 1) specify the arguments directly in the `tidy()` call
tidy(gravity, conf.int = TRUE, cluster = c("Product", "Year"))
tidy(gravity, conf.int = TRUE, se = "threeway")
# 2) or, feed tidy() a summary.fixest object that has already accepted
# these arguments
gravity_summ <- summary(gravity, cluster = c("Product", "Year"))</pre>
tidy(gravity_summ, conf.int = TRUE)
# approach (1) is preferred.
```

tidy.ftable

Description

This function is deprecated. Please use tibble::as_tibble() instead.

Usage

S3 method for class 'ftable'
tidy(x, ...)

Arguments

| х | An ftable object returned from <pre>stats::ftable().</pre> |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

An ftable contains a "flat" contingency table. This melts it into a tibble::tibble with one column for each variable, then a Freq column.

See Also

Other deprecated: bootstrap(), confint_tidy(), data.frame_tidiers, finish_glance(), fix_data_frame(), summary_tidiers, tidy.density(), tidy.dist(), tidy.numeric()

tidy.Gam

Tidy a(n) Gam object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'Gam'
tidy(x, ...)
```

Arguments

| х | A Gam object returned from a call to gam::gam(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

Tidy gam objects created by calls to mgcv::gam() with tidy.gam().

Value

A tibble::tibble() with columns:

| df | Degrees of freedom used by this term in the model. |
|-----------|---|
| meansq | Mean sum of squares. Equal to total sum of squares divided by degrees of freedom. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| sumsq | Sum of squares explained by this term. |
| term | The name of the regression term. |
| | |

See Also

tidy(), gam::gam(), tidy.anova(), tidy.gam() Other gam tidiers: glance.Gam()

Examples

load libraries for models and data library(gam)

fit model

tidy.gam

```
g <- gam(mpg ~ s(hp, 4) + am + qsec, data = mtcars)
# summarize model fit with tidiers
tidy(g)
glance(g)</pre>
```

tidy.gam

Tidy a(n) gam object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'gam'
tidy(
    x,
    parametric = FALSE,
    conf.int = FALSE,
    conf.level = 0.95,
    exponentiate = FALSE,
    ...
)
```

Arguments

| х | A gam object returned from a call to mgcv::gam(). |
|--------------|--|
| parametric | Logical indicating if parametric or smooth terms should be tidied. Defaults to FALSE, meaning that smooth terms are tidied by default. |
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

When parametric = FALSE return columns edf and ref.df rather than estimate and std.error.

Value

A tibble::tibble() with columns:

| estimate | The estimated value of the regression term. |
|-----------|---|
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| edf | The effective degrees of freedom. Only reported when 'parametric = FALSE' |
| ref.df | The reference degrees of freedom. Only reported when 'parametric = FALSE' |
| | |

See Also

tidy(), mgcv::gam()

Other mgcv tidiers: glance.gam()

```
# load libraries for models and data
library(mgcv)
# fit model
g <- gam(mpg ~ s(hp) + am + qsec, data = mtcars)
# summarize model fit with tidiers
tidy(g)
tidy(g, parametric = TRUE)
glance(g)
augment(g)</pre>
```

tidy.garch

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'garch'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| х | A garch object returned by tseries::garch(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- |
| | zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), tseries::garch()

Other garch tidiers: glance.garch()

Examples

```
# load libraries for models and data
library(tseries)
# load data
data(EuStockMarkets)
# fit model
dax <- diff(log(EuStockMarkets))[, "DAX"]
dax.garch <- garch(dax)
dax.garch
# summarize model fit with tidiers
tidy(dax.garch)</pre>
```

tidy.geeglm Tidy a(n) geeglm object

Description

glance(dax.garch)

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'geeglm'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)
```

Arguments

| х | A geeglm object returned from a call to geepack::geeglm(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |

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| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
|--------------|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| atoila | |

Details

If conf.int = TRUE, the confidence interval is computed with the an internal confint.geeglm() function.

If you have missing values in your model data, you may need to refit the model with na.action = na.exclude or deal with the missingness in the data beforehand.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), geepack::geeglm()

Examples

```
# load modeling library
library(geepack)
```

load data
data(state)

```
ds <- data.frame(state.region, state.x77)
# fit model
geefit <- geeglm(Income ~ Frost + Murder,
    id = state.region,
    data = ds,
    corstr = "exchangeable"
)
# summarize model fit with tidiers
tidy(geefit)
tidy(geefit, conf.int = TRUE)</pre>
```

tidy.glht

Tidy a(n) glht object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'glht'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| х | A glht object returned by multcomp::glht(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | - |

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tidy.glht

Value

A tibble::tibble() with columns:

| contrast | Levels being compared. |
|------------|---|
| estimate | The estimated value of the regression term. |
| null.value | Value to which the estimate is compared. |

See Also

tidy(), multcomp::glht()

Other multcomp tidiers: tidy.cld(), tidy.confint.glht(), tidy.summary.glht()

```
# load libraries for models and data
library(multcomp)
library(ggplot2)
amod <- aov(breaks ~ wool + tension, data = warpbreaks)</pre>
wht <- glht(amod, linfct = mcp(tension = "Tukey"))</pre>
tidy(wht)
ggplot(wht, aes(lhs, estimate)) +
  geom_point()
CI <- confint(wht)</pre>
tidy(CI)
ggplot(CI, aes(lhs, estimate, ymin = lwr, ymax = upr)) +
  geom_pointrange()
tidy(summary(wht))
ggplot(mapping = aes(lhs, estimate)) +
  geom_linerange(aes(ymin = lwr, ymax = upr), data = CI) +
  geom_point(aes(size = p), data = summary(wht)) +
  scale_size(trans = "reverse")
cld <- cld(wht)</pre>
tidy(cld)
```

tidy.glm

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'glm'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)
```

Arguments

| x | A glm object returned from stats::glm(). |
|--------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will |

See Also

stats::glm()

be ignored.

Other Im tidiers: augment.glm(), augment.lm(), glance.glm(), glance.lm(), glance.summary.lm(), glance.svyglm(), tidy.lm.beta(), tidy.lm(), tidy.mlm(), tidy.summary.lm() tidy.glmnet

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'glmnet'
tidy(x, return_zeros = FALSE, ...)
```

Arguments

| x return_zeros | A glmnet object returned from glmnet::glmnet(). Logical indicating whether coefficients with value zero zero should be included in the results. Defaults to FALSE. |
|-------------------|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

Note that while this representation of GLMs is much easier to plot and combine than the default structure, it is also much more memory-intensive. Do not use for large, sparse matrices.

No augment method is yet provided even though the model produces predictions, because the input data is not tidy (it is a matrix that may be very wide) and therefore combining predictions with it is not logical. Furthermore, predictions make sense only with a specific choice of lambda.

Value

A tibble::tibble() with columns:

| dev.ratio | Fraction of null deviance explained at each value of lambda. |
|-----------|--|
| estimate | The estimated value of the regression term. |
| lambda | Value of penalty parameter lambda. |
| step | Which step of lambda choices was used. |
| term | The name of the regression term. |

tidy.glmRob

See Also

tidy(), glmnet::glmnet()
Other glmnet tidiers: glance.cv.glmnet(), glance.glmnet(), tidy.cv.glmnet()

Examples

```
# load libraries for models and data
library(glmnet)
set.seed(2014)
x <- matrix(rnorm(100 * 20), 100, 20)
y <- rnorm(100)
fit1 <- glmnet(x, y)</pre>
# summarize model fit with tidiers + visualization
tidy(fit1)
glance(fit1)
library(dplyr)
library(ggplot2)
tidied <- tidy(fit1) %>% filter(term != "(Intercept)")
ggplot(tidied, aes(step, estimate, group = term)) +
 geom_line()
ggplot(tidied, aes(lambda, estimate, group = term)) +
 geom_line() +
 scale_x_log10()
ggplot(tidied, aes(lambda, dev.ratio)) +
 geom_line()
# works for other types of regressions as well, such as logistic
g2 <- sample(1:2, 100, replace = TRUE)
fit2 <- glmnet(x, g2, family = "binomial")</pre>
tidy(fit2)
```

tidy.glmRob

Tidy a(n) glmRob object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

tidy.glmRob

Usage

```
## S3 method for class 'glmRob'
tidy(x, ...)
```

Arguments

| x | A glmRob object returned from robust::glmRob(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

For tidiers for robust models from the MASS package see tidy.rlm().

See Also

robust::glmRob()

Other robust tidiers: augment.lmRob(), glance.glmRob(), glance.lmRob(), tidy.lmRob()

Examples

```
# load libraries for models and data
library(robust)
# fit model
gm <- glmRob(am ~ wt, data = mtcars, family = "binomial")</pre>
# summarize model fit with tidiers
tidy(gm)
glance(gm)
```

tidy.glmrob

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'glmrob'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| x | A glmrob object returned from robustbase::glmrob(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

For tidiers for robust models from the MASS package see tidy.rlm().

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |

tidy.gmm

| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- |
|-----------|--|
| | zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

robustbase::glmrob()

Other robustbase tidiers: augment.glmrob(), augment.lmrob(), glance.lmrob(), tidy.lmrob()

Examples

```
if (requireNamespace("robustbase", quietly = TRUE)) {
 # load libraries for models and data
 library(robustbase)
 data(coleman)
 set.seed(0)
 m <- lmrob(Y ~ ., data = coleman)</pre>
 tidy(m)
 augment(m)
 glance(m)
 data(carrots)
 Rfit <- glmrob(cbind(success, total - success) ~ logdose + block,</pre>
    family = binomial, data = carrots, method = "Mqle",
    control = glmrobMqle.control(tcc = 1.2)
 )
 tidy(Rfit)
 augment(Rfit)
}
```

tidy.gmm

Tidy a(n) gmm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'gmm'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)
```

Arguments

| x | A gmm object returned from gmm::gmm(). |
|--------------|--|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- |
| | zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), gmm::gmm()

Other gmm tidiers: glance.gmm()

Examples

load libraries for models and data
library(gmm)

examples come from the "gmm" package

tidy.gmm

```
# CAPM test with GMM
data(Finance)
r <- Finance[1:300, 1:10]
rm <- Finance[1:300, "rm"]</pre>
rf <- Finance[1:300, "rf"]</pre>
z <- as.matrix(r - rf)</pre>
t <- nrow(z)
zm <- rm - rf
h <- matrix(zm, t, 1)</pre>
res <- gmm(z \sim zm, x = h)
# tidy result
tidy(res)
tidy(res, conf.int = TRUE)
tidy(res, conf.int = TRUE, conf.level = .99)
# coefficient plot
library(ggplot2)
library(dplyr)
tidy(res, conf.int = TRUE) %>%
  mutate(variable = reorder(term, estimate)) %>%
  ggplot(aes(estimate, variable)) +
  geom_point() +
  geom_errorbarh(aes(xmin = conf.low, xmax = conf.high)) +
  geom_vline(xintercept = 0, color = "red", lty = 2)
# from a function instead of a matrix
g <- function(theta, x) {</pre>
  e <- x[, 2:11] - theta[1] - (x[, 1] - theta[1]) %*% matrix(theta[2:11], 1, 10)</pre>
  gmat <- cbind(e, e * c(x[, 1]))</pre>
  return(gmat)
}
x <- as.matrix(cbind(rm, r))</pre>
res_black <- gmm(g, x = x, t0 = rep(0, 11))
tidy(res_black)
tidy(res_black, conf.int = TRUE)
# APT test with Fama-French factors and GMM
f1 <- zm
f2 <- Finance[1:300, "hml"] - rf</pre>
f3 <- Finance[1:300, "smb"] - rf
h \leq cbind(f1, f2, f3)
res2 <- gmm(z \sim f1 + f2 + f3, x = h)
td2 <- tidy(res2, conf.int = TRUE)</pre>
td2
# coefficient plot
```

```
td2 %>%
  mutate(variable = reorder(term, estimate)) %>%
  ggplot(aes(estimate, variable)) +
  geom_point() +
  geom_errorbarh(aes(xmin = conf.low, xmax = conf.high)) +
  geom_vline(xintercept = 0, color = "red", lty = 2)
```

tidy.htest

Tidy/glance a(n) htest object

Description

For models that have only a single component, the tidy() and glance() methods are identical. Please see the documentation for both of those methods.

Usage

```
## S3 method for class 'htest'
tidy(x, ...)
```

S3 method for class 'htest'
glance(x, ...)

Arguments

| х | <pre>An htest objected, such as those created by stats::cor.test(), stats::t.test(), stats::wilcox.test(), stats::chisq.test(), etc.</pre> |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| alternative | Alternative hypothesis (character). |
|-------------|--|
| conf.high | Upper bound on the confidence interval for the estimate. |
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |

tidy.ivreg

| estimate1 | Sometimes two estimates are computed, such as in a two-sample t-test. |
|-----------|---|
| estimate2 | Sometimes two estimates are computed, such as in a two-sample t-test. |
| method | Method used. |
| p.value | The two-sided p-value associated with the observed statistic. |
| parameter | The parameter being modeled. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |

See Also

tidy(), stats::cor.test(), stats::t.test(), stats::wilcox.test(), stats::chisq.test()
Other htest tidiers: augment.htest(), tidy.pairwise.htest(), tidy.power.htest()

Examples

```
tt <- t.test(rnorm(10))
tidy(tt)
# the glance output will be the same for each of the below tests
glance(tt)
tt <- t.test(mpg ~ am, data = mtcars)
tidy(tt)
wt <- wilcox.test(mpg ~ am, data = mtcars, conf.int = TRUE, exact = FALSE)
tidy(wt)
ct <- cor.test(mtcars$wt, mtcars$mpg)
tidy(ct)
chit <- chisq.test(xtabs(Freq ~ Sex + Class, data = as.data.frame(Titanic)))
tidy(chit)
augment(chit)</pre>
```

tidy.ivreg

Tidy a(n) ivreg object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'ivreg'
tidy(x, conf.int = FALSE, conf.level = 0.95, instruments = FALSE, ...)
```

Arguments

| x | An ivreg object created by a call to AER::ivreg(). |
|-------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| instruments | Logical indicating whether to return coefficients from the second-stage or diag- nostics tests for each endogenous regressor (F-statistics). Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

This tidier currently only supports ivreg-classed objects outputted by the AER package. The ivreg package also outputs objects of class ivreg, and will be supported in a later release.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|----------------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| p.value.Sargan | p-value for Sargan test of overidentifying restrictions. |

tidy.ivreg

| p.value.weakins | it | |
|----------------------|--|--|
| | p-value for weak instruments test. | |
| p.value.Wu.Hausman | | |
| | p-value for Wu-Hausman weak instruments test for endogeneity. | |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- | |
| | zero. | |
| statistic.Sargan | | |
| | Statistic for Sargan test of overidentifying restrictions. | |
| statistic.weakinst | | |
| | Statistic for Wu-Hausman test. | |
| statistic.Wu.Hausman | | |
| | Statistic for Wu-Hausman weak instruments test for endogeneity. | |
| std.error | The standard error of the regression term. | |
| term | The name of the regression term. | |
| | | |

See Also

tidy(), AER::ivreg()

Other ivreg tidiers: augment.ivreg(), glance.ivreg()

Examples

```
# load libraries for models and data
library(AER)
# load data
data("CigarettesSW", package = "AER")
# fit model
ivr <- ivreg(</pre>
 log(packs) ~ income | population,
 data = CigarettesSW,
  subset = year == "1995"
)
# summarize model fit with tidiers
tidy(ivr)
tidy(ivr, conf.int = TRUE)
tidy(ivr, conf.int = TRUE, instruments = TRUE)
augment(ivr)
augment(ivr, data = CigarettesSW)
augment(ivr, newdata = CigarettesSW)
glance(ivr)
```

tidy.kappa

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'kappa'
tidy(x, ...)

Arguments

х

. . .

A kappa object returned from psych::cohen.kappa().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

Note that confidence level (alpha) for the confidence interval cannot be set in tidy. Instead you must set the alpha argument to psych::cohen.kappa() when creating the kappa object.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| type | Either 'weighted' or 'unweighted'. |

See Also

tidy(), psych::cohen.kappa()

tidy.kde

Examples

```
# load libraries for models and data
library(psych)
# generate example data
rater1 <- 1:9
rater2 <- c(1, 3, 1, 6, 1, 5, 5, 6, 7)
# fit model
ck <- cohen.kappa(cbind(rater1, rater2))
# summarize model fit with tidiers + visualization
tidy(ck)
# graph the confidence intervals
library(ggplot2)
ggplot(tidy(ck), aes(estimate, type)) +
geom_point() +
geom_errorbarh(aes(xmin = conf.low, xmax = conf.high))
```

tidy.kde

Tidy a(n) kde object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'kde'
tidy(x, ...)
```

Arguments

- x A kde object returned from ks::kde().
- ... Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in ..., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

```
Returns a data frame in long format with four columns. Use tidyr::pivot_wider(..., names_from = variable, values_from = value) on the output to return to a wide format.
```

Value

A tibble::tibble() with columns:

| estimate | The estimated value of the regression term. |
|----------|---|
| obs | weighted observed number of events in each group. |
| value | The value/estimate of the component. Results from data reshaping. |
| variable | Variable under consideration. |

See Also

tidy(), ks::kde()

Examples

```
# load libraries for models and data
library(ks)
# generate data
dat <- replicate(2, rnorm(100))</pre>
k \leq kde(dat)
# summarize model fit with tidiers + visualization
td <- tidy(k)</pre>
td
library(ggplot2)
library(dplyr)
library(tidyr)
td %>%
  pivot_wider(c(obs, estimate),
    names_from = variable,
    values_from = value
  ) %>%
  ggplot(aes(x1, x2, fill = estimate)) +
  geom_tile() +
  theme_void()
```

tidy.Kendall

```
# also works with 3 dimensions
dat3 <- replicate(3, rnorm(100))
k3 <- kde(dat3)
td3 <- tidy(k3)
td3</pre>
```

tidy.Kendall

Tidy a(n) Kendall object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'Kendall'
tidy(x, ...)

Arguments

| х | A Kendall object returned from a call to Kendall::Kendall(), Kendall::MannKendall(), or Kendall::SeasonalMannKendall(). |
|-----|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |
| lue | |

Value

A tibble::tibble() with columns:

| kendall_score | Kendall score. | |
|-------------------|---|--|
| p.value | The two-sided p-value associated with the observed statistic. | |
| var_kendall_score | | |
| | Variance of the kendall_score. | |
| statistic | Kendall's tau statistic | |
| denominator | The denominator, which is tau=kendall_score/denominator. | |

See Also

tidy(),Kendall::Kendall(),Kendall::MannKendall(),Kendall::SeasonalMannKendall()

Examples

```
# load libraries for models and data
library(Kendall)
A <- c(2.5, 2.5, 2.5, 2.5, 2.5, 5, 6.5, 6.5, 10, 10, 10, 10, 10, 14, 14, 14, 16, 17)
B <- c(1, 1, 1, 1, 2, 1, 1, 2, 1, 1, 1, 1, 1, 1, 2, 2, 2)
# fit models and summarize results
f_res <- Kendall(A, B)
tidy(f_res)
s_res <- MannKendall(B)
tidy(s_res)
t_res <- SeasonalMannKendall(ts(A))
tidy(t_res)
```

tidy.kmeans

Tidy a(n) kmeans object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'kmeans'
tidy(x, col.names = colnames(x$centers), ...)
```

Arguments

| х | A kmeans object created by stats::kmeans(). |
|-----------|---|
| col.names | Dimension names. Defaults to the names of the variables in x. Set to NULL to get names $x1, x2, \ldots$ |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |

tidy.lavaan

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| cluster | A factor describing the cluster from 1:k. |
|----------|---|
| size | Number of points assigned to cluster. |
| withinss | The within-cluster sum of squares. |

See Also

tidy(), stats::kmeans()

Other kmeans tidiers: augment.kmeans(), glance.kmeans()

Examples

```
library(cluster)
library(modeldata)
library(dplyr)
data(hpc_data)
x <- hpc_data[, 2:5]
fit <- pam(x, k = 4)
tidy(fit)
glance(fit)
augment(fit, x)</pre>
```

tidy.lavaan

Tidy a(n) lavaan object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'lavaan'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| x | A lavaan object, such as those returned from $lavaan::cfa()$, and $lavaan::sem()$. |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments passed to lavaan::parameterEstimates(). Caution- ary note: Misspecified arguments may be silently ignored. |

Value

A tibble::tibble() with one row for each estimated parameter and columns:

| term | The result of paste(lhs, op, rhs) |
|-----------|--|
| ор | The operator in the model syntax (e.g. ~~ for covariances, or ~ for regression parameters) |
| group | The group (if specified) in the lavaan model |
| estimate | The parameter estimate (may be standardized) |
| std.error | |
| statistic | The z value returned by lavaan::parameterEstimates() |
| p.value | |
| conf.low | |
| conf.high | |
| std.lv | Standardized estimates based on the variances of the (continuous) latent variables only |
| std.all | Standardized estimates based on both the variances of both (continuous) ob- served and latent variables. |
| std.nox | Standardized estimates based on both the variances of both (continuous) ob- served and latent variables, but not the variances of exogenous covariates. |

See Also

tidy(), lavaan::cfa(), lavaan::sem(), lavaan::parameterEstimates()
Other lavaan tidiers: glance.lavaan()

tidy.lm

Examples

```
# load libraries for models and data
library(lavaan)

cfa.fit <- cfa("F =~ x1 + x2 + x3 + x4 + x5 + x6 + x7 + x8 + x9",
    data = HolzingerSwineford1939, group = "school"
)
tidy(cfa.fit)</pre>
```

tidy.lm

Tidy a(n) lm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'lm'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)
```

Arguments

| x | An lm object created by stats::lm(). |
|--------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: • tidy() methods will warn when supplied an exponentiate argument if it |
| | will be ignored.augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

If the linear model is an mlm object (multiple linear model), there is an additional column response. See tidy.mlm().

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), stats::summary.lm()

Other Im tidiers: augment.glm(), augment.lm(), glance.glm(), glance.lm(), glance.summary.lm(), glance.svyglm(), tidy.glm(), tidy.lm.beta(), tidy.mlm(), tidy.summary.lm()

Examples

```
library(ggplot2)
library(dplyr)
mod <- lm(mpg \sim wt + qsec, data = mtcars)
tidy(mod)
glance(mod)
# coefficient plot
d <- tidy(mod, conf.int = TRUE)</pre>
ggplot(d, aes(estimate, term, xmin = conf.low, xmax = conf.high, height = 0)) +
  geom_point() +
  geom_vline(xintercept = 0, lty = 4) +
  geom_errorbarh()
# aside: There are tidy() and glance() methods for lm.summary objects too.
# this can be useful when you want to conserve memory by converting large lm
# objects into their leaner summary.lm equivalents.
s <- summary(mod)</pre>
tidy(s, conf.int = TRUE)
glance(s)
```

```
tidy.lm
```

```
augment(mod)
augment(mod, mtcars, interval = "confidence")
# predict on new data
newdata <- mtcars %>%
 head(6) %>%
 mutate(wt = wt + 1)
augment(mod, newdata = newdata)
# ggplot2 example where we also construct 95% prediction interval
# simpler bivariate model since we're plotting in 2D
mod2 <- lm(mpg ~ wt, data = mtcars)</pre>
au <- augment(mod2, newdata = newdata, interval = "prediction")</pre>
ggplot(au, aes(wt, mpg)) +
 geom_point() +
 geom_line(aes(y = .fitted)) +
 geom_ribbon(aes(ymin = .lower, ymax = .upper), col = NA, alpha = 0.3)
# predict on new data without outcome variable. Output does not include .resid
newdata <- newdata %>%
 select(-mpg)
augment(mod, newdata = newdata)
au <- augment(mod, data = mtcars)</pre>
ggplot(au, aes(.hat, .std.resid)) +
 geom_vline(size = 2, colour = "white", xintercept = 0) +
 geom_hline(size = 2, colour = "white", yintercept = 0) +
 geom_point() +
 geom_smooth(se = FALSE)
plot(mod, which = 6)
ggplot(au, aes(.hat, .cooksd)) +
 geom_vline(xintercept = 0, colour = NA) +
 geom_abline(slope = seq(0, 3, by = 0.5), colour = "white") +
 geom_smooth(se = FALSE) +
 geom_point()
# column-wise models
a <- matrix(rnorm(20), nrow = 10)</pre>
b <- a + rnorm(length(a))</pre>
result <- lm(b ~ a)</pre>
tidy(result)
```

tidy.lm.beta

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'lm.beta'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| х | An lm.beta object created by lm.beta::lm.beta. |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Details

If the linear model is an mlm object (multiple linear model), there is an additional column response.

If you have missing values in your model data, you may need to refit the model with na.action = na.exclude.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |

tidy.lmodel2

| estimate | The estimated value of the regression term. |
|-----------|---|
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

```
Other Im tidiers: augment.glm(), augment.lm(), glance.glm(), glance.lm(), glance.summary.lm(),
glance.svyglm(), tidy.glm(), tidy.lm(), tidy.mlm(), tidy.summary.lm()
```

Examples

```
# load libraries for models and data
library(lm.beta)
# fit models
mod <- stats::lm(speed ~ ., data = cars)</pre>
std <- lm.beta(mod)</pre>
# summarize model fit with tidiers
tidy(std, conf.int = TRUE)
# generate data
ctl <- c(4.17, 5.58, 5.18, 6.11, 4.50, 4.61, 5.17, 4.53, 5.33, 5.14)
trt <- c(4.81, 4.17, 4.41, 3.59, 5.87, 3.83, 6.03, 4.89, 4.32, 4.69)
group <- gl(2, 10, 20, labels = c("Ctl", "Trt"))</pre>
weight <- c(ctl, trt)</pre>
# fit models
mod2 <- lm(weight ~ group)</pre>
std2 <- lm.beta(mod2)</pre>
# summarize model fit with tidiers
tidy(std2, conf.int = TRUE)
```

tidy.lmodel2 Tidy a(n) lmodel2 object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'lmodel2'
tidy(x, ...)
```

Arguments

х

A lmodel2 object returned by lmodel2::lmodel2(). Additional arguments. Not used. Needed to match generic signature only. Cau-. . . tionary note: Misspelled arguments will be absorbed in ..., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

There are always only two terms in an Imodel2: "Intercept" and "Slope". These are computed by four methods: OLS (ordinary least squares), MA (major axis), SMA (standard major axis), and RMA (ranged major axis).

The returned p-value is one-tailed and calculated via a permutation test. A permutational test is used because distributional assumptions may not be valid. More information can be found in vignette("mod2user", package = "lmodel2").

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| term | The name of the regression term. |
| method | Either OLS/MA/SMA/RMA |

See Also

tidy(), lmodel2::lmodel2()

Other Imodel2 tidiers: glance.lmodel2()

tidy.lmRob

Examples

```
# load libraries for models and data
library(lmodel2)
data(mod2ex2)
Ex2.res <- lmodel2(Prey ~ Predators, data = mod2ex2, "relative", "relative", 99)
Ex2.res
# summarize model fit with tidiers + visualization
tidy(Ex2.res)
glance(Ex2.res)
# this allows coefficient plots with ggplot2
library(ggplot2)
ggplot(tidy(Ex2.res), aes(estimate, term, color = method)) +
geom_point() +
geom_errorbarh(aes(xmin = conf.low, xmax = conf.high)) +
geom_errorbarh(aes(xmin = conf.low, xmax = conf.high)) +
geom_errorbarh(aes(xmin = conf.low, xmax = conf.high))
```

tidy.lmRob

Tidy a(n) lmRob object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'lmRob'
tidy(x, ...)
```

Arguments

| Х | |
|---|--|
| | |
| | |

A lmRob object returned from robust::lmRob().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

• tidy() methods will warn when supplied an exponentiate argument if it will be ignored.

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

For tidiers for robust models from the MASS package see tidy.rlm().

See Also

robust::lmRob()

Other robust tidiers: augment.lmRob(), glance.glmRob(), glance.lmRob(), tidy.glmRob()

Examples

```
# load modeling library
library(robust)
# fit model
m <- lmRob(mpg ~ wt, data = mtcars)
# summarize model fit with tidiers
tidy(m)
augment(m)
glance(m)
```

tidy.lmrob

Tidy a(n) lmrob object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'lmrob'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

tidy.lmrob

Arguments

| x | A lmrob object returned from robustbase::lmrob(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

For tidiers for robust models from the MASS package see tidy.rlm().

See Also

robustbase::lmrob()

Other robustbase tidiers: augment.glmrob(), augment.lmrob(), glance.lmrob(), tidy.glmrob()

Examples

```
if (requireNamespace("robustbase", quietly = TRUE)) {
    # load libraries for models and data
    library(robustbase)
    data(coleman)
    set.seed(0)
    m <- lmrob(Y ~ ., data = coleman)
    tidy(m)
    augment(m)
    glance(m)
    data(carrots)
    Rfit <- glmrob(cbind(success, total - success) ~ logdose + block,
    family = binomial, data = carrots, method = "Mqle",
    control = glmrobMqle.control(tcc = 1.2)
    )
    tidy(Rfit)</pre>
```

```
augment(Rfit)
}
```

tidy.lsmobj

Tidy a(n) lsmobj object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'lsmobj'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| х | An 1smobj object. |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments passed to emmeans::summary.emmGrid() or lsmeans::summary.ref.grid(). Cautionary note : misspecified arguments may be silently ignored! |

Details

Returns a data frame with one observation for each estimated marginal mean, and one column for each combination of factors. When the input is a contrast, each row will contain one estimated contrast.

There are a large number of arguments that can be passed on to emmeans::summary.emmGrid() or lsmeans::summary.ref.grid().

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|------------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| contrast | Levels being compared. |
| df | Degrees of freedom used by this term in the model. |
| null.value | Value to which the estimate is compared. |

tidy.lsmobj

| p.value | The two-sided p-value associated with the observed statistic. |
|-----------|---|
| std.error | The standard error of the regression term. |
| estimate | Expected marginal mean |
| statistic | T-ratio statistic |

See Also

```
tidy(), emmeans::ref_grid(), emmeans::emmeans(), emmeans::contrast()
Other emmeans tidiers: tidy.emmGrid(), tidy.ref.grid(), tidy.summary_emm()
```

Examples

```
# load libraries for models and data
library(emmeans)
# linear model for sales of oranges per day
oranges_lm1 <- lm(sales1 ~ price1 + price2 + day + store, data = oranges)</pre>
# reference grid; see vignette("basics", package = "emmeans")
oranges_rg1 <- ref_grid(oranges_lm1)</pre>
td <- tidy(oranges_rg1)</pre>
td
# marginal averages
marginal <- emmeans(oranges_rg1, "day")</pre>
tidy(marginal)
# contrasts
tidy(contrast(marginal))
tidy(contrast(marginal, method = "pairwise"))
# plot confidence intervals
library(ggplot2)
ggplot(tidy(marginal, conf.int = TRUE), aes(day, estimate)) +
  geom_point() +
  geom_errorbar(aes(ymin = conf.low, ymax = conf.high))
# by multiple prices
by_price <- emmeans(oranges_lm1, "day",</pre>
 by = "price2",
  at = list(
    price1 = 50, price2 = c(40, 60, 80),
    day = c("2", "3", "4")
  )
)
by_price
```

```
tidy(by_price)
ggplot(tidy(by_price, conf.int = TRUE), aes(price2, estimate, color = day)) +
  geom_line() +
  geom_errorbar(aes(ymin = conf.low, ymax = conf.high))
# joint_tests
tidy(joint_tests(oranges_lm1))
```

tidy.manova Tia

Tidy a(n) manova object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'manova'
tidy(x, test = "Pillai", ...)

Arguments

| х | A manova object return from stats::manova(). |
|------|---|
| test | One of "Pillai" (Pillai's trace), "Wilks" (Wilk's lambda), "Hotelling-Lawley" (Hotelling-Lawley trace) or "Roy" (Roy's greatest root) indicating which test statistic should be used. Defaults to "Pillai". |
| | Arguments passed on to stats::summary.manova |
| | object An object of class "manova" or an aov object with multiple responses.intercept logical. If TRUE, the intercept term is included in the table.tol tolerance to be used in deciding if the residuals are rank-deficient: see qr. |

Details

Depending on which test statistic is specified only one of pillai, wilks, hl or roy is included.

Value

A tibble::tibble() with columns:

| den.df | Degrees of freedom of the denominator. |
|---------|---|
| num.df | Degrees of freedom. |
| p.value | The two-sided p-value associated with the observed statistic. |

tidy.map

| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- |
|-----------|--|
| | zero. |
| term | The name of the regression term. |
| pillai | Pillai's trace. |
| wilks | Wilk's lambda. |
| hl | Hotelling-Lawley trace. |
| roy | Roy's greatest root. |

See Also

```
tidy(), stats::summary.manova()
```

```
Other anova tidiers: glance.anova(), glance.aov(), tidy.TukeyHSD(), tidy.anova(), tidy.aovlist(), tidy.aov()
```

Examples

npk2 <- within(npk, foo <- rnorm(24))
m <- manova(cbind(yield, foo) ~ block + N * P * K, npk2)
tidy(m)</pre>

```
tidy.map
```

Tidy a(n) map object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'map'
tidy(x, ...)
```

Arguments

| x | A map object returned from maps::map(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

tidy.margins

Value

| A tibble::tibble() with columns: | | |
|----------------------------------|--|--|
| rm. | | |
| | | |
| | | |
| | | |

Remaining columns give information on geographic attributes and depend on the inputted map object. See ?maps::map for more information.

See Also

tidy(), maps::map()

Examples

```
# load libraries for models and data
library(maps)
library(ggplot2)
ca <- map("county", "ca", plot = FALSE, fill = TRUE)
tidy(ca)
qplot(long, lat, data = ca, geom = "polygon", group = group)
tx <- map("county", "texas", plot = FALSE, fill = TRUE)
tidy(tx)
qplot(long, lat,
    data = tx, geom = "polygon", group = group,
    colour = I("white")
)
```

tidy.margins Tidy a(n) margins object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'margins'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

tidy.margins

Arguments

| х | A margins object returned from margins::margins(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

The margins package provides a way to obtain coefficient marginal effects for a variety of (nonlinear) models, such as logit or models with multiway interaction terms. Note that the glance.margins() method requires rerunning the underlying model again, which can take some time. Similarly, an augment.margins() method is not currently supported, but users can simply run the underlying model to obtain the same information.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), margins::margins()

tidy.Mclust

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'Mclust' tidy(x, ...)

Arguments

| x | An Mclust object return from mclust::Mclust(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| proportion | The mixing proportion of each component |
|------------|---|
| size | Number of points assigned to cluster. |
| mean | The mean for each component. In case of 2+ dimensional models, a column with the mean is added for each dimension. NA for noise component |
| variance | In case of one-dimensional and spherical models, the variance for each compo- nent, omitted otherwise. NA for noise component |
| component | Cluster id as a factor. |

See Also

tidy(), mclust::Mclust()

Other mclust tidiers: augment.Mclust()

tidy.mediate

Examples

```
# load library for models and data
library(mclust)
# load data manipulation libraries
library(dplyr)
library(tibble)
library(purrr)
library(tidyr)
set.seed(27)
centers <- tibble(</pre>
  cluster = factor(1:3),
  # number points in each cluster
  num_points = c(100, 150, 50),
  # x1 coordinate of cluster center
  x1 = c(5, 0, -3),
  # x2 coordinate of cluster center
  x^2 = c(-1, 1, -2)
)
points <- centers %>%
  mutate(
    x1 = map2(num_points, x1, rnorm),
    x2 = map2(num_points, x2, rnorm)
  ) %>%
  select(-num_points, -cluster) %>%
  unnest(c(x1, x2))
# fit model
m <- Mclust(points)</pre>
# summarize model fit with tidiers
tidy(m)
augment(m, points)
glance(m)
```

tidy.mediate

Tidy a(n) mediate object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'mediate'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| х | A mediate object produced by a call to mediation::mediate(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

The tibble has four rows. The first two indicate the mediated effect in the control and treatment groups, respectively. And the last two the direct effect in each group.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), mediation::mediate()

tidy.mfx

Examples

```
# load libraries for models and data
library(mediation)
data(jobs)
# fit models
b <- lm(job_seek ~ treat + econ_hard + sex + age, data = jobs)
c <- lm(depress2 ~ treat + job_seek + econ_hard + sex + age, data = jobs)
mod <- mediate(b, c, sims = 50, treat = "treat", mediator = "job_seek")
# summarize model fit with tidiers
tidy(mod)
tidy(mod, conf.int = TRUE)
tidy(mod, conf.int = TRUE, conf.level = .99)
```

```
tidy.mfx
```

Tidy a(n) mfx object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

The particular functions below provide generic tidy methods for objects returned by the mfx package, preserving the calculated marginal effects instead of the naive model coefficients. The returned tidy tibble will also include an additional "atmean" column indicating how the marginal effects were originally calculated (see Details below).

Usage

```
## S3 method for class 'mfx'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
## S3 method for class 'logitmfx'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
## S3 method for class 'negbinmfx'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
## S3 method for class 'poissonmfx'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
## S3 method for class 'probitmfx'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| x | A logitmfx, negbinmfx, poissonmfx, or probitmfx object. (Note that betamfx objects receive their own set of tidiers.) |
|------------|--|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

The mfx package provides methods for calculating marginal effects for various generalized linear models (GLMs). Unlike standard linear models, estimated model coefficients in a GLM cannot be directly interpreted as marginal effects (i.e., the change in the response variable predicted after a one unit change in one of the regressors). This is because the estimated coefficients are multiplicative, dependent on both the link function that was used for the estimation and any other variables that were included in the model. When calculating marginal effects, users must typically choose whether they want to use i) the average observation in the data, or ii) the average of the sample marginal effects. See vignette("mfxarticle") from the mfx package for more details.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| atmean | TRUE if the marginal effects were originally calculated as the partial effects for the average observation. If FALSE, then these were instead calculated as average partial effects. |

tidy.mjoint

See Also

```
tidy(), mfx::logitmfx(), mfx::negbinmfx(), mfx::poissonmfx(), mfx::probitmfx()
Other mfx tidiers: augment.betamfx(), augment.mfx(), glance.betamfx(), glance.mfx(),
```

Examples

tidy.betamfx()

```
# load libraries for models and data
library(mfx)
# get the marginal effects from a logit regression
mod_logmfx <- logitmfx(am ~ cyl + hp + wt, atmean = TRUE, data = mtcars)</pre>
tidy(mod_logmfx, conf.int = TRUE)
# compare with the naive model coefficients of the same logit call
tidy(
 glm(am ~ cyl + hp + wt, family = binomial, data = mtcars),
 conf.int = TRUE
)
augment(mod_logmfx)
glance(mod_logmfx)
# another example, this time using probit regression
mod_probmfx <- probitmfx(am ~ cyl + hp + wt, atmean = TRUE, data = mtcars)</pre>
tidy(mod_probmfx, conf.int = TRUE)
augment(mod_probmfx)
glance(mod_probmfx)
```

tidy.mjoint Tidy a(n) mjoint object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'mjoint'
tidy(
```

```
x,
component = "survival",
conf.int = FALSE,
conf.level = 0.95,
boot_se = NULL,
...
```

Arguments

| х | An mjoint object returned from joineRML::mjoint(). |
|------------|--|
| component | Character specifying whether to tidy the survival or the longitudinal component of the model. Must be either "survival" or "longitudinal". Defaults to "survival". |
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| boot_se | Optionally a bootSE object from joineRML::bootSE(). If specified, calculates confidence intervals via the bootstrap. Defaults to NULL, in which case standard errors are calculated from the empirical information matrix. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

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tidy.mjoint

See Also

tidy(), joineRML::mjoint(), joineRML::bootSE()
Other mjoint tidiers: glance.mjoint()

Examples

```
# broom only skips running these examples because the example models take a
# while to generatethey should run just fine, though!
## Not run:
# load libraries for models and data
library(joineRML)
# fit a joint model with bivariate longitudinal outcomes
data(heart.valve)
hvd <- heart.valve[!is.na(heart.valve$log.grad) &</pre>
 !is.na(heart.valve$log.lvmi) &
 heart.valve$num <= 50, ]</pre>
fit <- mjoint(</pre>
 formLongFixed = list(
    "grad" = log.grad ~ time + sex + hs,
    "lvmi" = log.lvmi ~ time + sex
 ).
 formLongRandom = list(
    "grad" = \sim 1 | num,
    "lvmi" = ~ time | num
 ),
 formSurv = Surv(fuyrs, status) ~ age,
 data = hvd,
 inits = list("gamma" = c(0.11, 1.51, 0.80)),
 timeVar = "time"
)
# extract the survival fixed effects
tidy(fit)
# extract the longitudinal fixed effects
tidy(fit, component = "longitudinal")
# extract the survival fixed effects with confidence intervals
tidy(fit, ci = TRUE)
# extract the survival fixed effects with confidence intervals based
# on bootstrapped standard errors
bSE <- bootSE(fit, nboot = 5, safe.boot = TRUE)</pre>
tidy(fit, boot_se = bSE, ci = TRUE)
```

```
# augment original data with fitted longitudinal values and residuals
hvd2 <- augment(fit)
# extract model statistics
glance(fit)
## End(Not run)</pre>
```

tidy.mle2

Tidy a(n) mle2 object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'mle2'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| х | An mle2 object created by a call to bbmle::mle2(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

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tidy.mlm

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), bbmle::mle2(), tidy_optim()

Examples

```
# load libraries for models and data
library(bbmle)
# generate data
x <- 0:10
y <- c(26, 17, 13, 12, 20, 5, 9, 8, 5, 4, 8)
d <- data.frame(x, y)
# fit model
fit <- mle2(y ~ dpois(lambda = ymean),
   start = list(ymean = mean(y)), data = d
)
# summarize model fit with tidiers
tidy(fit)</pre>
```

tidy.mlm

Tidy a(n) mlm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'mlm'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| х | An mlm object created by stats::lm() with a matrix as the response. |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

In contrast to lm object (simple linear model), tidy output for mlm (multiple linear model) objects contain an additional column response.

If you have missing values in your model data, you may need to refit the model with na.action = na.exclude.

Value

A tibble::tibble() with columns:

| Upper bound on the confidence interval for the estimate. |
|--|
| Lower bound on the confidence interval for the estimate. |
| The estimated value of the regression term. |
| The two-sided p-value associated with the observed statistic. |
| The value of a T-statistic to use in a hypothesis that the regression term is non- |
| zero. |
| The standard error of the regression term. |
| The name of the regression term. |
| |

See Also

tidy()

Other Im tidiers: augment.glm(), augment.lm(), glance.glm(), glance.lm(), glance.summary.lm(), glance.svyglm(), tidy.glm(), tidy.lm.beta(), tidy.lm(), tidy.summary.lm()

tidy.mlogit

Examples

```
# fit model
mod <- lm(cbind(mpg, disp) ~ wt, mtcars)
# summarize model fit with tidiers
tidy(mod, conf.int = TRUE)</pre>
```

tidy.mlogit Tidying methods for logit models

Description

These methods tidy the coefficients of mnl and nl models generated by the functions of the mlogit package.

Usage

```
## S3 method for class 'mlogit'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| х | an object returned from mlogit::mlogit(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |

| estimate | The estimated value of the regression term. |
|-----------|---|
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), mlogit::mlogit()

Other mlogit tidiers: augment.mlogit(), glance.mlogit()

Examples

```
# load libraries for models and data
library(mlogit)
data("Fishing", package = "mlogit")
Fish <- dfidx(Fishing, varying = 2:9, shape = "wide", choice = "mode")
# fit model
m <- mlogit(mode ~ price + catch | income, data = Fish)
# summarize model fit with tidiers
tidy(m)
augment(m)
glance(m)
```

tidy.muhaz

Tidy a(n) muhaz object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'muhaz'
tidy(x, ...)
```

tidy.multinom

Arguments

| х | A muhaz object returned by muhaz::muhaz(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| time | Point in time. |
|----------|------------------------|
| estimate | Estimated hazard rate. |

See Also

tidy(), muhaz::muhaz()

Other muhaz tidiers: glance.muhaz()

Examples

```
# load libraries for models and data
library(muhaz)
library(survival)
# fit model
x <- muhaz(ovarian$futime, ovarian$fustat)</pre>
```

```
# summarize model fit with tidiers
tidy(x)
glance(x)
```

tidy.multinom Tidying methods for multinomial logistic regression models

Description

These methods tidy the coefficients of multinomial logistic regression models generated by multinom of the nnet package.

Usage

```
## S3 method for class 'multinom'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)
```

Arguments

| х | A multinom object returned from nnet::multinom(). |
|--------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | 6 |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| y.value | The response level. |

See Also

tidy(), nnet::multinom()

Other multinom tidiers: glance.multinom()

tidy.negbin

Examples

```
# load libraries for models and data
library(nnet)
library(MASS)
example(birthwt)
bwt.mu <- multinom(low ~ ., bwt)
tidy(bwt.mu)
glance(bwt.mu)
# or, for output from a multinomial logistic regression
fit.gear <- multinom(gear ~ mpg + factor(am), data = mtcars)
tidy(fit.gear)
glance(fit.gear)
```

tidy.negbin

Tidy a(n) negbin object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'negbin'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)
```

Arguments

| х | A glm.nb object returned by MASS::glm.nb(). |
|--------------|--|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |

| Additional arguments. Not used. Needed to match generic signature only. Cau- |
|--|
| tionary note: Misspelled arguments will be absorbed in, where they will be |
| ignored. If the misspelled argument has a default value, the default value will be |
| used. For example, if you pass conf.lvel = 0.9, all computation will proceed |
| using conf.level = 0.95. Two exceptions here are: |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

See Also

MASS::glm.nb()

Other glm.nb tidiers: glance.negbin()

Examples

```
# load libraries for models and data
library(MASS)
# fit model
r <- glm.nb(Days ~ Sex / (Age + Eth * Lrn), data = quine)
# summarize model fit with tidiers
tidy(r)
glance(r)</pre>
```

tidy.nlrq

Tidy a(n) nlrq object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'nlrq'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

. . .

tidy.nlrq

Arguments

| x | A nlrq object returned from quantreg::nlrq(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), quantreg::nlrq()

Other quantreg tidiers: augment.nlrq(), augment.rqs(), augment.rq(), glance.nlrq(), glance.rq(), tidy.rqs(), tidy.rq()

Examples

```
# load modeling library
library(quantreg)
# build artificial data with multiplicative error
set.seed(1)
dat <- NULL</pre>
```

tidy.nls

```
dat$x <- rep(1:25, 20)
dat$y <- SSlogis(dat$x, 10, 12, 2) * rnorm(500, 1, 0.1)
# fit the median using nlrq
mod <- nlrq(y ~ SSlogis(x, Asym, mid, scal),
    data = dat, tau = 0.5, trace = TRUE
)
# summarize model fit with tidiers
tidy(mod)
glance(mod)
augment(mod)</pre>
```

tidy.nls

Tidy a(n) nls object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'nls'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| x | An nls object returned from stats::nls(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

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tidy.nls

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

```
tidy, stats::nls(), stats::summary.nls()
```

Other nls tidiers: augment.nls(), glance.nls()

Examples

```
# fit model
n <- nls(mpg ~ k * e^wt, data = mtcars, start = list(k = 1, e = 2))
# summarize model fit with tidiers + visualization
tidy(n)
augment(n)
glance(n)
library(ggplot2)
ggplot(augment(n), aes(wt, mpg)) +
geom_point() +
geom_line(aes(y = .fitted))
newdata <- head(mtcars)
newdata$wt <- newdata$wt + 1
augment(n, newdata = newdata)
```

tidy.numeric

Description

Vector tidiers are deprecated and will be removed from an upcoming release of broom.

Usage

```
## S3 method for class 'numeric'
tidy(x, ...)
## S3 method for class 'character'
tidy(x, ...)
## S3 method for class 'logical'
tidy(x, ...)
```

Arguments

| Х | An object of class "numeric", "integer", "character", or "logical". Most likely a |
|---|---|
| | named vector |
| | Extra arguments (not used) |

Details

Turn atomic vectors into data frames, where the names of the vector (if they exist) are a column and the values of the vector are a column.

See Also

Other deprecated: bootstrap(), confint_tidy(), data.frame_tidiers, finish_glance(), fix_data_frame(), summary_tidiers, tidy.density(), tidy.dist(), tidy.ftable()

Other deprecated: bootstrap(), confint_tidy(), data.frame_tidiers, finish_glance(), fix_data_frame(), summary_tidiers, tidy.density(), tidy.dist(), tidy.ftable()

Other deprecated: bootstrap(), confint_tidy(), data.frame_tidiers, finish_glance(), fix_data_frame(), summary_tidiers, tidy.density(), tidy.dist(), tidy.ftable()

Examples

```
## Not run:
x <- 1:5
names(x) <- letters[1:5]
tidy(x)
```

End(Not run)

tidy.orcutt

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'orcutt'
tidy(x, ...)
```

Arguments

| An orcutt object returned from orcutt::cochrane.orcutt(). |
|--|
| Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are: |
| tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will |
| |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| estimate | The estimated value of the regression term. |
|-----------|---|
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

orcutt::cochrane.orcutt()

Other orcutt tidiers: glance.orcutt()

Examples

```
# load libraries for models and data
library(orcutt)
# fit model and summarize results
reg <- lm(mpg ~ wt + qsec + disp, mtcars)
tidy(reg)
co <- cochrane.orcutt(reg)
tidy(co)
glance(co)
```

tidy.pairwise.htest Tidy a(n) pairwise.htest object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'pairwise.htest'
tidy(x, ...)
```

Arguments

| х | A pairwise.htest object such as those returned from stats::pairwise.t.test() or stats::pairwise.wilcox.test(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

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tidy.pam

Details

Note that in one-sided tests, the alternative hypothesis of each test can be stated as "group1 is greater/less than group2".

Note also that the columns of group1 and group2 will always be a factor, even if the original input is (e.g.) numeric.

Value

A tibble::tibble() with columns:

| group1 | First group being compared. |
|---------|---|
| group2 | Second group being compared. |
| p.value | The two-sided p-value associated with the observed statistic. |

See Also

stats::pairwise.t.test(), stats::pairwise.wilcox.test(), tidy()
Other htest tidiers: augment.htest(), tidy.htest(), tidy.power.htest()

Examples

```
attach(airquality)
Month <- factor(Month, labels = month.abb[5:9])
ptt <- pairwise.t.test(Ozone, Month)
tidy(ptt)
library(modeldata)
data(hpc_data)
attach(hpc_data)
ptt2 <- pairwise.t.test(compounds, class)
tidy(ptt2)
tidy(pairwise.t.test(compounds, class, alternative = "greater"))
tidy(pairwise.t.test(compounds, class, alternative = "less"))
tidy(pairwise.wilcox.test(compounds, class))
```

tidy.pam

Tidy a(n) pam object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'pam'
tidy(x, col.names = paste0("x", 1:ncol(x$medoids)), ...)
```

Arguments

| x | An pam object returned from cluster::pam() |
|-----------|---|
| col.names | Column names in the input data frame. Defaults to the names of the variables in x. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

For examples, see the pam vignette.

Value

A tibble::tibble() with columns:

| size | Size of each cluster. |
|------------|--|
| max.diss | Maximal dissimilarity between the observations in the cluster and that cluster's medoid. |
| avg.diss | Average dissimilarity between the observations in the cluster and that cluster's medoid. |
| diameter | Diameter of the cluster. |
| separation | Separation of the cluster. |
| avg.width | Average silhouette width of the cluster. |
| cluster | A factor describing the cluster from 1:k. |

See Also

tidy(), cluster::pam()

Other pam tidiers: augment.pam(), glance.pam()

tidy.plm

Examples

```
# load libraries for models and data
library(dplyr)
library(ggplot2)
library(cluster)
library(modeldata)
data(hpc_data)
x <- hpc_data[, 2:5]</pre>
p <- pam(x, k = 4)
# summarize model fit with tidiers + visualization
tidy(p)
glance(p)
augment(p, x)
augment(p, x) %>%
  ggplot(aes(compounds, input_fields)) +
  geom_point(aes(color = .cluster)) +
  geom_text(aes(label = cluster), data = tidy(p), size = 10)
```

Tidy a(n) plm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'plm'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| x | A plm objected returned by plm::plm(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |

| Additional arguments. Not used. Needed to match generic signature only. Cau- |
|--|
| tionary note: Misspelled arguments will be absorbed in, where they will be |
| ignored. If the misspelled argument has a default value, the default value will be |
| used. For example, if you pass conf.lvel = 0.9, all computation will proceed |
| using conf.level = 0.95. Two exceptions here are: |

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| | |

See Also

tidy(), plm::plm(), tidy.lm()

Other plm tidiers: augment.plm(), glance.plm()

Examples

```
# load libraries for models and data
library(plm)
# load data
data("Produc", package = "plm")
# fit model
zz <- plm(log(gsp) ~ log(pcap) + log(pc) + log(emp) + unemp,
data = Produc, index = c("state", "year")
)
# summarize model fit with tidiers
summary(zz)
tidy(zz)
tidy(zz, conf.int = TRUE)
```

. . .

tidy.poLCA

```
tidy(zz, conf.int = TRUE, conf.level = 0.9)
augment(zz)
glance(zz)
```

```
tidy.poLCA
```

Tidy a(n) poLCA object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'poLCA'
tidy(x, ...)
```

Arguments

| x | A poLCA object returned from poLCA::poLCA(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with columns:

| class | The class under consideration. |
|-----------|--|
| outcome | Outcome of manifest variable. |
| std.error | The standard error of the regression term. |
| variable | Manifest variable |
| estimate | Estimated class-conditional response probability |
| | |

See Also

tidy(), poLCA::poLCA()
Other poLCA tidiers: augment.poLCA(), glance.poLCA()

Examples

```
# load libraries for models and data
library(poLCA)
library(dplyr)
# generate data
data(values)
f <- cbind(A, B, C, D) ~ 1
# fit model
M1 <- poLCA(f, values, nclass = 2, verbose = FALSE)
M1
# summarize model fit with tidiers + visualization
tidy(M1)
augment(M1)
glance(M1)
library(ggplot2)
ggplot(tidy(M1), aes(factor(class), estimate, fill = factor(outcome))) +
  geom_bar(stat = "identity", width = 1) +
  facet_wrap(~variable)
# three-class model with a single covariate.
data(election)
f2a <- cbind(
  MORALG, CARESG, KNOWG, LEADG, DISHONG, INTELG,
  MORALB, CARESB, KNOWB, LEADB, DISHONB, INTELB
) ~ PARTY
nes2a <- poLCA(f2a, election, nclass = 3, nrep = 5, verbose = FALSE)</pre>
td <- tidy(nes2a)</pre>
td
ggplot(td, aes(outcome, estimate, color = factor(class), group = class)) +
  geom_line() +
  facet_wrap(~variable, nrow = 2) +
  theme(axis.text.x = element_text(angle = 90, hjust = 1))
au <- augment(nes2a)</pre>
```

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tidy.polr

```
au
count(au, .class)
# if the original data is provided, it leads to NAs in new columns
# for rows that weren't predicted
au2 <- augment(nes2a, data = election)
au2
dim(au2)
```

tidy.polr *Tidy a(n) polr object*

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'polr'
tidy(
    x,
    conf.int = FALSE,
    conf.level = 0.95,
    exponentiate = FALSE,
    p.values = FALSE,
    ...
)
```

Arguments

| x | A polr object returned from MASS::polr(). |
|--------------|--|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |

| p.values | Logical. Should p-values be returned, based on chi-squared tests from MASS::dropterm(). Defaults to FALSE. |
|----------|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will |
| | be ignored. |

Details

In broom 0.7.0 the coefficient_type column was renamed to coef.type, and the contents were changed as well. Now the contents are coefficient and scale, rather than coefficient and zeta.

Calculating p.values with the dropterm() function is the approach suggested by the MASS package author. This approach is computationally intensive so that p.values are only returned if requested explicitly. Additionally, it only works for models containing no variables with more than two categories. If this condition is not met, a message is shown and NA is returned instead of p-values.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy, MASS::polr()

Other ordinal tidiers: augment.clm(), augment.polr(), glance.clmm(), glance.clm(), glance.polr(), glance.svyolr(), tidy.clmm(), tidy.clm(), tidy.svyolr()

Examples

load libraries for models and data
library(MASS)

```
# fit model
fit <- polr(Sat ~ Infl + Type + Cont, weights = Freq, data = housing)
# summarize model fit with tidiers
tidy(fit, exponentiate = TRUE, conf.int = TRUE)
glance(fit)
augment(fit, type.predict = "class")
fit2 <- polr(factor(gear) ~ am + mpg + qsec, data = mtcars)
tidy(fit, p.values = TRUE)
```

tidy.power.htest Tidy a(n) power.htest object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'power.htest'
tidy(x, ...)

Arguments ×

. . .

A power.htest object such as those returned from stats::power.t.test().

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . ., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

delta True difference in means.

| n | Number of observations by component. |
|-----------|--|
| power | Power achieved for given value of n. |
| sd | Standard deviation. |
| sig.level | Significance level (Type I error probability). |

See Also

```
stats::power.t.test()
```

```
Other htest tidiers: augment.htest(), tidy.htest(), tidy.pairwise.htest()
```

Examples

```
ptt <- power.t.test(n = 2:30, delta = 1)
tidy(ptt)
library(ggplot2)
ggplot(tidy(ptt), aes(n, power)) +
  geom_line()</pre>
```

tidy.prcomp Tidy a(n) prcomp object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'prcomp'
tidy(x, matrix = "u", ...)
```

Arguments

| х | A prcomp object returned by stats::prcomp(). |
|--------|---|
| matrix | Character specifying which component of the PCA should be tidied. |
| | • "u", "samples", "scores", or "x": returns information about the map from the original space into principle components space. |
| | • "v", "rotation", "loadings" or "variables": returns information about the map from principle components space back into the original space. |

. . .

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . ., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

See https://stats.stackexchange.com/questions/134282/relationship-between-svd-and-pca-how-to-usesvd-to-perform-pca for information on how to interpret the various tidied matrices. Note that SVD is only equivalent to PCA on centered data.

Value

A tibble::tibble with columns depending on the component of PCA being tidied.

If matrix is "u", "samples", "scores", or "x" each row in the tidied output corresponds to the original data in PCA space. The columns are:

| row | ID of the original observation (i.e. rowname from original data). |
|-------|--|
| PC | Integer indicating a principal component. |
| value | The score of the observation for that particular principal component. That is, the location of the observation in PCA space. |

If matrix is "v", "rotation", "loadings" or "variables", each row in the tidied output corresponds to information about the principle components in the original space. The columns are:

| row | The variable labels (colnames) of the data set on which PCA was performed. |
|--------------------|--|
| PC | An integer vector indicating the principal component. |
| value | The value of the eigenvector (axis score) on the indicated principal component. |
| If matrix is "d",' | 'eigenvalues" or "pcs", the columns are: |
| PC | An integer vector indicating the principal component. |
| std.dev | Standard deviation explained by this PC. |
| percent | Fraction of variation explained by this component (a numeric value between 0 and 1). |
| cumulative | Cumulative fraction of variation explained by principle components up to this component (a numeric value between 0 and 1). |

See Also

stats::prcomp(), svd_tidiers

Other svd tidiers: augment.prcomp(), tidy_irlba(), tidy_svd()

Examples

```
pc <- prcomp(USArrests, scale = TRUE)</pre>
# information about rotation
tidy(pc)
# information about samples (states)
tidy(pc, "samples")
# information about PCs
tidy(pc, "pcs")
# state map
library(dplyr)
library(ggplot2)
library(maps)
pc %>%
  tidy(matrix = "samples") %>%
  mutate(region = tolower(row)) %>%
  inner_join(map_data("state"), by = "region") %>%
  ggplot(aes(long, lat, group = group, fill = value)) +
  geom_polygon() +
  facet_wrap(~PC) +
  theme_void() +
  ggtitle("Principal components of arrest data")
au <- augment(pc, data = USArrests)</pre>
au
ggplot(au, aes(.fittedPC1, .fittedPC2)) +
  geom_point() +
  geom_text(aes(label = .rownames), vjust = 1, hjust = 1)
```

tidy.pyears

Tidy a(n) pyears object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

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tidy.pyears

Usage

```
## S3 method for class 'pyears'
tidy(x, ...)
```

Arguments

| х | A pyears object returned from survival::pyears(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

expected is only present in the output when if a ratetable term is present.

If the data.frame = TRUE argument is supplied to pyears, this is simply the contents of x\$data.

Value

A tibble::tibble() with columns:

| expected | Expected number of events. |
|----------|--------------------------------------|
| pyears | Person-years of exposure. |
| n | number of subjects contributing time |
| event | observed number of events |

See Also

tidy(), survival::pyears()

Other pyears tidiers: glance.pyears()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(),
glance.coxph(),glance.pyears(),glance.survdiff(),glance.survexp(),glance.survfit(),
glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.survdiff(), tidy.survexp(),
tidy.survfit(), tidy.survreg()
```

Examples

load libraries for models and data library(survival)

```
# generate and format data
temp.yr <- tcut(mgus$dxyr, 55:92, labels = as.character(55:91))</pre>
temp.age <- tcut(mgus$age, 34:101, labels = as.character(34:100))</pre>
ptime <- ifelse(is.na(mgus$pctime), mgus$futime, mgus$pctime)</pre>
pstat <- ifelse(is.na(mgus$pctime), 0, 1)</pre>
pfit <- pyears(Surv(ptime / 365.25, pstat) ~ temp.yr + temp.age + sex, mgus,</pre>
  data.frame = TRUE
)
# summarize model fit with tidiers
tidy(pfit)
glance(pfit)
# if data.frame argument is not given, different information is present in
# output
pfit2 <- pyears(Surv(ptime / 365.25, pstat) ~ temp.yr + temp.age + sex, mgus)</pre>
tidy(pfit2)
glance(pfit2)
```

tidy.rcorr

Tidy a(n) rcorr object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'rcorr'
tidy(x, diagonal = FALSE, ...)

Arguments

| х | An rcorr object returned from Hmisc::rcorr(). |
|----------|---|
| diagonal | Logical indicating whether or not to include diagonal elements of the correlation matrix, or the correlation of a column with itself. For the elements, estimate is always 1 and p.value is always NA. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |

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tidy.rcorr

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

Suppose the original data has columns A and B. In the correlation matrix from rcorr there may be entries for both the cor(A, B) and cor(B, A). Only one of these pairs will ever be present in the tidy output.

Value

A tibble::tibble() with columns:

| column1 | Name or index of the first column being described. |
|----------|---|
| column2 | Name or index of the second column being described. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| n | Number of observations used to compute the correlation |

See Also

tidy(), Hmisc::rcorr()

Examples

```
# load libraries for models and data
library(Hmisc)
mat <- replicate(52, rnorm(100))
# add some NAs
mat[sample(length(mat), 2000)] <- NA
# also, column names
colnames(mat) <- c(LETTERS, letters)
# fit model
rc <- rcorr(mat)
# summarize model fit with tidiers + visualization
td <- tidy(rc)
td
library(ggplot2)
ggplot(td, aes(p.value)) +
geom_histogram(binwidth = .1)
```

```
ggplot(td, aes(estimate, p.value)) +
geom_point() +
scale_y_log10()
```

tidy.ref.grid Tidy a(n) ref.grid object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'ref.grid'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| x | A ref.grid object created by emmeans::ref_grid(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments passed to emmeans::summary.emmGrid() or lsmeans::summary.ref.grid(). Cautionary note : misspecified arguments may be silently ignored! |

Details

Returns a data frame with one observation for each estimated marginal mean, and one column for each combination of factors. When the input is a contrast, each row will contain one estimated contrast.

There are a large number of arguments that can be passed on to emmeans::summary.emmGrid() or lsmeans::summary.ref.grid().

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |

tidy.ref.grid

| df | Degrees of freedom used by this term in the model. |
|-----------|---|
| p.value | The two-sided p-value associated with the observed statistic. |
| std.error | The standard error of the regression term. |
| estimate | Expected marginal mean |
| statistic | T-ratio statistic |

See Also

```
tidy(), emmeans::ref_grid(), emmeans::emmeans(), emmeans::contrast()
Other emmeans tidiers: tidy.emmGrid(), tidy.lsmobj(), tidy.summary_emm()
```

Examples

```
# load libraries for models and data
library(emmeans)
# linear model for sales of oranges per day
oranges_lm1 <- lm(sales1 ~ price1 + price2 + day + store, data = oranges)</pre>
# reference grid; see vignette("basics", package = "emmeans")
oranges_rg1 <- ref_grid(oranges_lm1)</pre>
td <- tidy(oranges_rg1)</pre>
td
# marginal averages
marginal <- emmeans(oranges_rg1, "day")</pre>
tidy(marginal)
# contrasts
tidy(contrast(marginal))
tidy(contrast(marginal, method = "pairwise"))
# plot confidence intervals
library(ggplot2)
ggplot(tidy(marginal, conf.int = TRUE), aes(day, estimate)) +
  geom_point() +
  geom_errorbar(aes(ymin = conf.low, ymax = conf.high))
# by multiple prices
by_price <- emmeans(oranges_lm1, "day",</pre>
  by = "price2",
  at = list(
    price1 = 50, price2 = c(40, 60, 80),
    day = c("2", "3", "4")
 )
)
by_price
```

```
tidy(by_price)
ggplot(tidy(by_price, conf.int = TRUE), aes(price2, estimate, color = day)) +
  geom_line() +
  geom_errorbar(aes(ymin = conf.low, ymax = conf.high))
# joint_tests
tidy(joint_tests(oranges_lm1))
```

tidy.regsubsets Tidy a(n) regsubsets object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'regsubsets'
tidy(x, ...)

Arguments

| х | A regsubsets object created by leaps::regsubsets(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| r.squared | R squared statistic, or the percent of variation explained by the model |
|---------------|---|
| adj.r.squared | Adjusted R squared statistic |
| BIC | Bayesian information criterion for the component. |
| mallows_cp | Mallow's Cp statistic. |

tidy.ridgelm

See Also

tidy(), leaps::regsubsets()

Examples

load libraries for models and data
library(leaps)

fit model
all_fits <- regsubsets(hp ~ ., mtcars)</pre>

summarize model fit with tidiers
tidy(all_fits)

tidy.ridgelm

Tidy a(n) ridgelm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'ridgelm'
tidy(x, ...)
```

Arguments

| х | A ridgelm object returned from MASS::lm.ridge(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will |

Value

A tibble::tibble() with columns:

| GCV | Generalized cross validation error estimate. |
|----------|--|
| lambda | Value of penalty parameter lambda. |
| term | The name of the regression term. |
| estimate | estimate of scaled coefficient using this lambda |
| scale | Scaling factor of estimated coefficient |
| | |

See Also

tidy(), MASS::lm.ridge()

Other ridgelm tidiers: glance.ridgelm()

Examples

```
# load libraries for models and data
library(MASS)
names(longley)[1] <- "y"</pre>
# fit model and summarizd results
fit1 <- lm.ridge(y ~ ., longley)</pre>
tidy(fit1)
fit2 <- lm.ridge(y ~ ., longley, lambda = seq(0.001, .05, .001))</pre>
td2 <- tidy(fit2)
g2 <- glance(fit2)
# coefficient plot
library(ggplot2)
ggplot(td2, aes(lambda, estimate, color = term)) +
  geom_line()
# GCV plot
ggplot(td2, aes(lambda, GCV)) +
  geom_line()
# add line for the GCV minimizing estimate
ggplot(td2, aes(lambda, GCV)) +
  geom_line() +
  geom_vline(xintercept = g2$lambdaGCV, col = "red", lty = 2)
```

tidy.rlm

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'rlm'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| x | An rlm object returned by MASS::rlm(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

See Also

MASS::rlm()

Other rlm tidiers: augment.rlm(), glance.rlm()

tidy.rma

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'rma'
tidy(
    x,
    conf.int = FALSE,
    conf.level = 0.95,
    exponentiate = FALSE,
    include_studies = FALSE,
    measure = "GEN",
    ...
)
```

Arguments

| х | An rma object such as those created by metafor::rma(), metafor::rma.uni(), metafor::rma.glmm(), metafor::rma.mv(), or metafor::rma.peto(). | |
|-----------------|---|--|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. | |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. | |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. | |
| include_studies | | |
| | Logical. Should individual studies be included in the output? Defaults to FALSE. | |
| measure | Measure type. See metafor::escalc() | |
| | Additional arguments. Not used. Needed to match generic signature only. Cau- tionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: | |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. | |

tidy.roc

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- |
| | zero. |
| std.error | The standard error of the regression term. |
| term | The name of the individual study |
| type | The estimate type (summary vs individual study) |
| | |

Examples

```
# load libraries for models and data
library(metafor)
df <-
    escalc(
        measure = "RR",
        ai = tpos,
        bi = tneg,
        ci = cpos,
        di = cneg,
        data = dat.bcg
    )
meta_analysis <- rma(yi, vi, data = df, method = "EB")
tidy(meta_analysis)
```

tidy.roc

Tidy a(n) roc object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

tidy.roc

Usage

```
## S3 method for class 'roc'
tidy(x, ...)
```

Arguments

| x | An roc object returned from a call to AUC::roc(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| cutoff | The cutoff used for classification. Observations with predicted probabilities above this value were assigned class 1, and observations with predicted probabilities below this value were assigned class 0. |
|--------|---|
| fpr | False positive rate. |
| tpr | The true positive rate at the given cutoff. |

See Also

tidy(), AUC::roc()

Examples

```
# load libraries for models and data
library(AUC)
# load data
data(churn)
# fit model
r <- roc(churn$predictions, churn$labels)</pre>
# summarize with tidiers + visualization
td <- tidy(r)
td
```

library(ggplot2)

tidy.rq

```
ggplot(td, aes(fpr, tpr)) +
  geom_line()
# compare the ROC curves for two prediction algorithms
library(dplyr)
library(tidyr)
rocs <- churn %>%
  pivot_longer(contains("predictions"),
    names_to = "algorithm",
    values_to = "value"
    ) %>%
  nest(data = -algorithm) %>%
  mutate(tidy_roc = purrr::map(data, ~ tidy(roc(.x$value, .x$labels)))) %>%
  unnest(tidy_roc)
ggplot(rocs, aes(fpr, tpr, color = algorithm)) +
  geom_line()
```

tidy.rq

Tidy a(n) rq object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'rq'
tidy(x, se.type = NULL, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| x | An rq object returned from quantreg::rq(). |
|------------|--|
| se.type | Character specifying the method to use to calculate standard errors. Passed to quantreg::summary.rq() se argument. Defaults to "rank" if the sample size is less than 1000, otherwise defaults to "nid". |
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments passed to quantreg::summary.rq(). |

Details

If se.type = "rank" confidence intervals are calculated by summary.rq and statistic and p.value values are not returned. When only a single predictor is included in the model, no confidence intervals are calculated and the confidence limits are set to NA.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| | |

See Also

tidy(), quantreg::rq()

```
Other quantreg tidiers: augment.nlrq(), augment.rqs(), augment.rq(), glance.nlrq(), glance.rq(),
tidy.nlrq(), tidy.rqs()
```

Examples

```
# load modeling library and data
library(quantreg)
data(stackloss)
# median (l1) regression fit for the stackloss data.
mod1 <- rq(stack.loss ~ stack.x, .5)
# weighted sample median
mod2 <- rq(rnorm(50) ~ 1, weights = runif(50))
# summarize model fit with tidiers
tidy(mod1)
glance(mod1)
augment(mod2)
augment(mod2)
```

varying tau to generate an rqs object

tidy.rqs

```
mod3 <- rq(stack.loss ~ stack.x, tau = c(.25, .5))
tidy(mod3)
augment(mod3)
# glance cannot handle rqs objects like `mod3`--use a purrr
# `map`-based workflow instead</pre>
```

tidy.rqs

Tidy a(n) rqs object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'rqs'
tidy(x, se.type = "rank", conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| Х | An rqs object returned from quantreg::rq(). |
|------------|---|
| se.type | Character specifying the method to use to calculate standard errors. Passed to quantreg::summary.rq() se argument. Defaults to "rank". |
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments passed to quantreg::summary.rqs() |

Details

If se.type = "rank" confidence intervals are calculated by summary.rq. When only a single predictor is included in the model, no confidence intervals are calculated and the confidence limits are set to NA.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |
| quantile | Linear conditional quantile. |

See Also

```
tidy(), quantreg::rq()
```

```
Other quantreg tidiers: augment.nlrq(), augment.rqs(), augment.rq(), glance.nlrq(), glance.rq(),
tidy.nlrq(), tidy.rq()
```

Examples

```
# load modeling library and data
library(quantreg)
data(stackloss)
# median (11) regression fit for the stackloss data.
mod1 <- rq(stack.loss ~ stack.x, .5)</pre>
# weighted sample median
mod2 <- rq(rnorm(50) ~ 1, weights = runif(50))</pre>
# summarize model fit with tidiers
tidy(mod1)
glance(mod1)
augment(mod1)
tidy(mod2)
glance(mod2)
augment(mod2)
# varying tau to generate an rqs object
mod3 <- rq(stack.loss ~ stack.x, tau = c(.25, .5))</pre>
tidy(mod3)
augment(mod3)
```

tidy.sarlm

```
# glance cannot handle rqs objects like `mod3`--use a purrr
# `man` haard walflew instead
```

```
# `map`-based workflow instead
```

```
tidy.sarlm
```

Tidying methods for spatially autoregressive models

Description

These methods tidy the coefficients of spatial autoregression models generated by functions in the spatialreg package.

Usage

S3 method for class 'sarlm'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| х | An object returned from spatialreg::lagsarlm() or spatialreg::errorsarlm(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment () methods will warn when supplied a newdata argument if it will |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- |
| | zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), spatialreg::lagsarlm(), spatialreg::errorsarlm(), spatialreg::sacsarlm()
Other spatialreg tidiers: augment.sarlm(), glance.sarlm()

Examples

```
# load libraries for models and data
library(spatialreg)
library(spdep)
# load data
data(oldcol, package = "spdep")
listw <- nb2listw(COL.nb, style = "W")</pre>
# fit model
crime_sar <-</pre>
  lagsarlm(CRIME ~ INC + HOVAL,
    data = COL.OLD,
    listw = listw,
    method = "eigen"
  )
# summarize model fit with tidiers
tidy(crime_sar)
tidy(crime_sar, conf.int = TRUE)
glance(crime_sar)
augment(crime_sar)
# fit another model
crime_sem <- errorsarlm(CRIME ~ INC + HOVAL, data = COL.OLD, listw)</pre>
# summarize model fit with tidiers
tidy(crime_sem)
tidy(crime_sem, conf.int = TRUE)
glance(crime_sem)
augment(crime_sem)
# fit another model
crime_sac <- sacsarlm(CRIME ~ INC + HOVAL, data = COL.OLD, listw)</pre>
# summarize model fit with tidiers
tidy(crime_sac)
tidy(crime_sac, conf.int = TRUE)
glance(crime_sac)
augment(crime_sac)
```

tidy.spec

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'spec'
tidy(x, ...)
```

Arguments

| х | A spec object created by stats::spectrum(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will |

Value

A tibble::tibble() with columns:

| freq | Vector of frequencies at which the spectral density is estimated. |
|------|---|
| spec | Vector (for univariate series) or matrix (for multivariate series) of estimates of the spectral density at frequencies corresponding to freq. |

See Also

tidy(), stats::spectrum()

Other time series tidiers: tidy.acf(), tidy.ts(), tidy.zoo()

be ignored.

Examples

```
spc <- spectrum(lh)
tidy(spc)
library(ggplot2)
ggplot(tidy(spc), aes(freq, spec)) +
  geom_line()</pre>
```

tidy.speedglm Tidy a(n) speedglm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'speedglm'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)
```

Arguments

| х | A speedglm object returned from speedglm::speedglm(). |
|--------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

tidy.speedlm

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

speedglm::speedglm()

Other speedlm tidiers: augment.speedlm(), glance.speedglm(), glance.speedlm(), tidy.speedlm()

Examples

```
# load libraries for models and data
library(speedglm)
# generate data
clotting <- data.frame(
    u = c(5, 10, 15, 20, 30, 40, 60, 80, 100),
    lot1 = c(118, 58, 42, 35, 27, 25, 21, 19, 18)
)
# fit model
fit <- speedglm(lot1 ~ log(u), data = clotting, family = Gamma(log))
# summarize model fit with tidiers
tidy(fit)
glance(fit)</pre>
```

tidy.speedlm Tidy a(n) speedlm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'speedlm'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| A speedlm object returned from speedglm::speedlm(). |
|--|
| Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| augment() methods will warn when supplied a newdata argument if it will be ignored. |
| |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

speedglm::speedlm(),tidy.lm()

Other speedlm tidiers: augment.speedlm(), glance.speedglm(), glance.speedlm(), tidy.speedglm()

Examples

load modeling library
library(speedglm)

tidy.summary.glht

```
# fit model
mod <- speedlm(mpg ~ wt + qsec, data = mtcars, fitted = TRUE)
# summarize model fit with tidiers
tidy(mod)
glance(mod)
augment(mod)</pre>
```

tidy.summary.glht Tidy a(n) summary.glht object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'summary.glht'
tidy(x, ...)

Arguments

| Х | A summary.glht object created by calling multcomp::summary.glht() on a glht object created with multcomp::glht(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| contrast | Levels being compared. |
|------------|--|
| estimate | The estimated value of the regression term. |
| null.value | Value to which the estimate is compared. |
| p.value | The two-sided p-value associated with the observed statistic |

| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- |
|-----------|--|
| | zero. |
| std.error | The standard error of the regression term. |

See Also

```
tidy(), multcomp::summary.glht(), multcomp::glht()
Other multcomp tidiers: tidy.cld(), tidy.confint.glht(), tidy.glht()
```

Examples

```
# load libraries for models and data
library(multcomp)
library(ggplot2)
amod <- aov(breaks ~ wool + tension, data = warpbreaks)</pre>
wht <- glht(amod, linfct = mcp(tension = "Tukey"))</pre>
tidy(wht)
ggplot(wht, aes(lhs, estimate)) +
  geom_point()
CI <- confint(wht)</pre>
tidy(CI)
ggplot(CI, aes(lhs, estimate, ymin = lwr, ymax = upr)) +
  geom_pointrange()
tidy(summary(wht))
ggplot(mapping = aes(lhs, estimate)) +
  geom_linerange(aes(ymin = lwr, ymax = upr), data = CI) +
  geom_point(aes(size = p), data = summary(wht)) +
  scale_size(trans = "reverse")
cld <- cld(wht)</pre>
tidy(cld)
```

tidy.summary.lm Tidy a(n) summary.lm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

tidy.summary.lm

Usage

```
## S3 method for class 'summary.lm'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

| х | A summary.lm object created by stats::summary.lm(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

The tidy.summary.lm() method is a potentially useful alternative to tidy.lm(). For instance, if users have already converted large lm objects into their leaner summary.lm equivalents to conserve memory.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|--|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- |
| | zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), stats::summary.lm()

Other Im tidiers: augment.glm(), augment.lm(), glance.glm(), glance.lm(), glance.summary.lm(), glance.svyglm(), tidy.glm(), tidy.lm.beta(), tidy.lm(), tidy.mlm()

Examples

```
# fit model
mod <- lm(mpg ~ wt + qsec, data = mtcars)
modsumm <- summary(mod)
# summarize model fit with tidiers
tidy(mod, conf.int = TRUE)
# equivalent to the above
tidy(modsumm, conf.int = TRUE)
glance(mod)
# mostly the same, except for a few missing columns
glance(modsumm)</pre>
```

tidy.summary_emm Tidy a(n) summary_emm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'summary_emm'
tidy(x, null.value = NULL, ...)

Arguments

| х | A summary_emm object. |
|------------|---|
| null.value | Value to which estimate is compared. |
| | Additional arguments passed to emmeans::summary.emmGrid() or lsmeans::summary.ref.grid(). Cautionary note : misspecified arguments may be silently ignored! |

Details

Returns a data frame with one observation for each estimated marginal mean, and one column for each combination of factors. When the input is a contrast, each row will contain one estimated contrast.

There are a large number of arguments that can be passed on to emmeans::summary.emmGrid() or lsmeans::summary.ref.grid().

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|------------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| contrast | Levels being compared. |
| den.df | Degrees of freedom of the denominator. |
| df | Degrees of freedom used by this term in the model. |
| null.value | Value to which the estimate is compared. |
| num.df | Degrees of freedom. |
| p.value | The two-sided p-value associated with the observed statistic. |
| std.error | The standard error of the regression term. |
| level1 | One level of the factor being contrasted |
| level2 | The other level of the factor being contrasted |
| term | Model term in joint tests |
| estimate | Expected marginal mean |
| statistic | T-ratio statistic or F-ratio statistic |

See Also

tidy(), emmeans::ref_grid(), emmeans::emmeans(), emmeans::contrast()
Other emmeans tidiers: tidy.emmGrid(), tidy.lsmobj(), tidy.ref.grid()

Examples

```
# load libraries for models and data
library(emmeans)
# linear model for sales of oranges per day
oranges_lm1 <- lm(sales1 ~ price1 + price2 + day + store, data = oranges)
# reference grid; see vignette("basics", package = "emmeans")
oranges_rg1 <- ref_grid(oranges_lm1)
td <- tidy(oranges_rg1)
td
# marginal averages
marginal <- emmeans(oranges_rg1, "day")
tidy(marginal)
# contrasts
tidy(contrast(marginal))
tidy(contrast(marginal, method = "pairwise"))
```

```
# plot confidence intervals
library(ggplot2)
ggplot(tidy(marginal, conf.int = TRUE), aes(day, estimate)) +
  geom_point() +
  geom_errorbar(aes(ymin = conf.low, ymax = conf.high))
# by multiple prices
by_price <- emmeans(oranges_lm1, "day",</pre>
  by = "price2",
  at = list(
   price1 = 50, price2 = c(40, 60, 80),
    day = c("2", "3", "4")
  )
)
by_price
tidy(by_price)
ggplot(tidy(by_price, conf.int = TRUE), aes(price2, estimate, color = day)) +
  geom_line() +
  geom_errorbar(aes(ymin = conf.low, ymax = conf.high))
# joint_tests
tidy(joint_tests(oranges_lm1))
```

tidy.survdiff *Tidy a(n) survdiff object*

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'survdiff'
tidy(x, ...)
```

Arguments

| x | An survdiff object returned from survival::survdiff(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature of |

. Additional arguments. Not used. Needed to match generic signature only. **Cau-tionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be

used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| exp | Weighted expected number of events in each group. |
|-----|---|
| Ν | Number of subjects in each group. |
| obs | weighted observed number of events in each group. |

See Also

tidy(), survival::survdiff()

Other survdiff tidiers: glance.survdiff()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(), glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(), glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survexp(), tidy.survexp(), tidy.survfit(), tidy.survreg()
```

Examples

```
# load libraries for models and data
library(survival)
# fit model
s <- survdiff(
  Surv(time, status) ~ pat.karno + strata(inst),
  data = lung
)
# summarize model fit with tidiers
tidy(s)
glance(s)
```

tidy.survexp

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'survexp'
tidy(x, ...)

Arguments

| Additional arguments. Not used. Needed to match generic signature of | |
|---|----------------------------|
| tionary note: Misspelled arguments will be absorbed in , where the ignored. If the misspelled argument has a default value, the default value will be absorbed. For example, if you pass conf.lvel = 0.9 , all computation will using conf.level = 0.95 . Two exceptions here are: | hey will be lue will be |
| tidy() methods will warn when supplied an exponentiate arg will be ignored. | ument if it |

 augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| n.risk | Number of individuals at risk at time zero. |
|----------|---|
| time | Point in time. |
| estimate | Estimate survival |

See Also

tidy(), survival::survexp()

Other survexp tidiers: glance.survexp()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(), glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(), glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(), tidy.survfit(), tidy.survreg()
```

tidy.survfit

Examples

```
# load libraries for models and data
library(survival)
# fit model
sexpfit <- survexp(</pre>
  futime \sim 1,
  rmap = list(
    sex = "male",
    year = accept.dt,
    age = (accept.dt - birth.dt)
  ),
  method = "conditional",
  data = jasa
)
# summarize model fit with tidiers
tidy(sexpfit)
glance(sexpfit)
```

tidy.survfit *Tidy a(n) survfit object*

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'survfit'
tidy(x, ...)
```

Arguments

```
Х
```

An survfit object returned from survival::survfit().

```
Additional arguments. Not used. Needed to match generic signature only. Cau-
tionary note: Misspelled arguments will be absorbed in . . . , where they will be
ignored. If the misspelled argument has a default value, the default value will be
used. For example, if you pass conf.lvel = 0.9, all computation will proceed
using conf.level = 0.95. Two exceptions here are:
```

• tidy() methods will warn when supplied an exponentiate argument if it will be ignored.

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| n.censor | Number of censored events. |
| n.event | Number of events at time t. |
| n.risk | Number of individuals at risk at time zero. |
| std.error | The standard error of the regression term. |
| time | Point in time. |
| estimate | estimate of survival or cumulative incidence rate when multistate |
| state | state if multistate survfit object input |
| strata | strata if stratified survfit object input |

See Also

tidy(), survival::survfit()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(),
glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(),
glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(),
tidy.survexp(), tidy.survreg()
```

Examples

```
# load libraries for models and data
library(survival)
# fit model
cfit <- coxph(Surv(time, status) ~ age + sex, lung)
sfit <- survfit(cfit)
# summarize model fit with tidiers + visualization
tidy(sfit)
glance(sfit)
library(ggplot2)
ggplot(tidy(sfit), aes(time, estimate)) +
geom_line() +
geom_ribbon(aes(ymin = conf.low, ymax = conf.high), alpha = .25)
# multi-state
```

tidy.survreg

```
fitCI <- survfit(Surv(stop, status * as.numeric(event), type = "mstate") ~ 1,
    data = mgus1, subset = (start == 0)
)
td_multi <- tidy(fitCI)
td_multi
ggplot(td_multi, aes(time, estimate, group = state)) +
    geom_line(aes(color = state)) +
    geom_ribbon(aes(ymin = conf.low, ymax = conf.high), alpha = .25)
```

tidy.survreg *Tidy a(n) survreg object*

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'survreg'
tidy(x, conf.level = 0.95, conf.int = FALSE, ...)

Arguments

| х | An survreg object returned from survival::survreg(). |
|------------|---|
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), survival::survreg()

Other survreg tidiers: augment.survreg(), glance.survreg()

```
Other survival tidiers: augment.coxph(), augment.survreg(), glance.aareg(), glance.cch(),
glance.coxph(), glance.pyears(), glance.survdiff(), glance.survexp(), glance.survfit(),
glance.survreg(), tidy.aareg(), tidy.cch(), tidy.coxph(), tidy.pyears(), tidy.survdiff(),
tidy.survexp(), tidy.survfit()
```

Examples

```
# load libraries for models and data
library(survival)
# fit model
sr <- survreg(</pre>
  Surv(futime, fustat) ~ ecog.ps + rx,
 ovarian,
 dist = "exponential"
)
# summarize model fit with tidiers + visualization
tidy(sr)
augment(sr, ovarian)
glance(sr)
# coefficient plot
td <- tidy(sr, conf.int = TRUE)</pre>
library(ggplot2)
ggplot(td, aes(estimate, term)) +
  geom_point() +
  geom_errorbarh(aes(xmin = conf.low, xmax = conf.high), height = 0) +
```

tidy.svyglm

```
geom_vline(xintercept = 0)
```

tidy.svyglm

Tidy a(n) svyglm object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'svyglm'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)

Arguments

| х | A svyglm object returned from survey::svyglm(). |
|--------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |

See Also

survey::svyglm(), stats::glm()

tidy.svyolr

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'svyolr'
tidy(x, conf.int = FALSE, conf.level = 0.95, exponentiate = FALSE, ...)

Arguments

| x | A svyolr object returned from survey::svyolr(). |
|--------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| exponentiate | Logical indicating whether or not to exponentiate the the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

The tidy.svyolr() tidier is a light wrapper around tidy.polr(). However, the implementation for p-value calculation in tidy.polr() is both computationally intensive and specific to that model, so the p.values argument to tidy.svyolr() is currently ignored, and will raise a warning when passed.

tidy.systemfit

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non-zero. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy, survey::svyolr()

```
Other ordinal tidiers: augment.clm(), augment.polr(), glance.clmm(), glance.clm(), glance.polr(),
glance.svyolr(), tidy.clmm(), tidy.clm(), tidy.polr()
```

Examples

```
library(broom)
library(survey)

data(api)
dclus1 <- svydesign(id = ~dnum, weights = ~pw, data = apiclus1, fpc = ~fpc)
dclus1 <- update(dclus1, mealcat = cut(meals, c(0, 25, 50, 75, 100)))

m <- svyolr(mealcat ~ avg.ed + mobility + stype, design = dclus1)

m
tidy(m, conf.int = TRUE)</pre>
```

tidy.systemfit Tidy a(n) systemfit object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'systemfit'
tidy(x, conf.int = TRUE, conf.level = 0.95, ...)
```

Arguments

| x | A systemfit object produced by a call to systemfit::systemfit(). |
|------------|---|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | 1 1 1 1 1 1 1 1 1 1 |

• augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

This tidy method works with any model objects of class systemfit. Default returns a tibble of six columns.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. |
|-----------|---|
| conf.low | Lower bound on the confidence interval for the estimate. |
| estimate | The estimated value of the regression term. |
| p.value | The two-sided p-value associated with the observed statistic. |
| std.error | The standard error of the regression term. |
| term | The name of the regression term. |

See Also

tidy(), systemfit::systemfit()

tidy.table

Examples

```
set.seed(27)
# load libraries for models and data
library(systemfit)
# generate data
df <- data.frame(</pre>
 X = rnorm(100),
 Y = rnorm(100),
 Z = rnorm(100),
 W = rnorm(100)
)
# fit model
fit <- systemfit(formula = list(Y ~ Z, W ~ X), data = df, method = "SUR")</pre>
# summarize model fit with tidiers
tidy(fit)
tidy(fit, conf.int = TRUE)
```

tidy.table

Tidy a(n) table object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Deprecated. Please use tibble::as_tibble() instead.

Usage

```
## S3 method for class 'table'
tidy(x, ...)
```

Arguments

- х
- . . .

A base::table object.

Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in ..., where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

Details

Directly calls tibble::as_tibble() on a base::table object.

Value

A tibble::tibble in long-form containing frequency information for the table in a Freq column. The result is much like what you get from tidyr::pivot_longer().

See Also

tibble::as_tibble.table()

tidy.ts

Tidy a(n) ts object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'ts'
tidy(x, ...)
```

Arguments

| x | A univariate or multivariate ts times series object. |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

series column is only present for multivariate ts objects.

Value

A tibble::tibble() with columns:

| index | Index (i.e. date or time) for a 'ts' or 'zoo' object. | |
|--------|---|--|
| series | Name of the series (present only for multivariate time series). | |
| value | The value/estimate of the component. Results from data reshaping. | |

See Also

tidy(), stats::ts()

Other time series tidiers: tidy.acf(), tidy.spec(), tidy.zoo()

Examples

```
set.seed(678)
```

```
tidy(ts(1:10, frequency = 4, start = c(1959, 2)))
z <- ts(matrix(rnorm(300), 100, 3), start = c(1961, 1), frequency = 12)
colnames(z) <- c("Aa", "Bb", "Cc")
tidy(z)</pre>
```

tidy.TukeyHSD Tidy a(n) TukeyHSD object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

```
## S3 method for class 'TukeyHSD'
tidy(x, ...)
```

Arguments

| х | A TukeyHSD object return from <pre>stats::TukeyHSD().</pre> |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Value

A tibble::tibble() with columns:

| adj.p.value | P-value adjusted for multiple comparisons. | |
|---|--|--|
| conf.high | Upper bound on the confidence interval for the estimate. | |
| conf.low | Lower bound on the confidence interval for the estimate. | |
| contrast | Levels being compared. | |
| estimate | e The estimated value of the regression term. | |
| null.value Value to which the estimate is compared. | | |
| term | The name of the regression term. | |

See Also

tidy(), stats::TukeyHSD()

```
Other anova tidiers: glance.anova(), glance.aov(), tidy.anova(), tidy.aovlist(), tidy.aov(),
tidy.manova()
```

Examples

```
fm1 <- aov(breaks ~ wool + tension, data = warpbreaks)
thsd <- TukeyHSD(fm1, "tension", ordered = TRUE)
tidy(thsd)
# may include comparisons on multiple terms</pre>
```

```
fm2 <- aov(mpg ~ as.factor(gear) * as.factor(cyl), data = mtcars)
tidy(TukeyHSD(fm2))</pre>
```

tidy.varest

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

Usage

S3 method for class 'varest'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

Arguments

| x | A varest object produced by a call to vars::VAR(). | |
|------------|---|--|
| conf.int | Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE. | |
| conf.level | The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. | |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: | |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. | |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. | |

Details

The tibble has one row for each term in the regression. The component column indicates whether a particular term was used to model either the "mean" or "precision". Here the precision is the inverse of the variance, often referred to as phi. At least one term will have been used to model the precision phi.

The vars package does not include a confint method and does not report confidence intervals for varest objects. Setting the tidy argument conf.int = TRUE will return a warning.

Value

A tibble::tibble() with columns:

| conf.high | Upper bound on the confidence interval for the estimate. | |
|-----------|---|--|
| conf.low | Lower bound on the confidence interval for the estimate. | |
| estimate | The estimated value of the regression term. | |
| p.value | The two-sided p-value associated with the observed statistic. | |
| statistic | The value of a T-statistic to use in a hypothesis that the regression term is non- | |
| | zero. | |
| std.error | The standard error of the regression term. | |
| term | The name of the regression term. | |
| component | Whether a particular term was used to model the mean or the precision in the regression. See details. | |

See Also

tidy(), vars::VAR()

Examples

```
# load libraries for models and data
library(vars)
# load data
data("Canada", package = "vars")
# fit models
mod <- VAR(Canada, p = 1, type = "both")
# summarize model fit with tidiers
tidy(mod)
glance(mod)</pre>
```

tidy.zoo

Tidy a(n) zoo object

Description

Tidy summarizes information about the components of a model. A model component might be a single term in a regression, a single hypothesis, a cluster, or a class. Exactly what tidy considers to be a model component varies across models but is usually self-evident. If a model has several distinct types of components, you will need to specify which components to return.

tidy.zoo

Usage

```
## S3 method for class 'zoo'
tidy(x, ...)
```

Arguments

| х | A zoo object such as those created by zoo::zoo(). |
|---|---|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |

Value

A tibble::tibble() with columns:

| index | Index (i.e. date or time) for a 'ts' or 'zoo' object. |
|--------|---|
| series | Name of the series (present only for multivariate time series). |
| value | The value/estimate of the component. Results from data reshaping. |

See Also

tidy(), zoo::zoo()

Other time series tidiers: tidy.acf(), tidy.spec(), tidy.ts()

Examples

```
# load libraries for models and data
library(zoo)
library(ggplot2)
set.seed(1071)
# generate data
Z.index <- as.Date(sample(12450:12500, 10))
Z.data <- matrix(rnorm(30), ncol = 3)
colnames(Z.data) <- c("Aa", "Bb", "Cc")
Z <- zoo(Z.data, Z.index)
# summarize model fit with tidiers + visualization
tidy(Z)
```

```
ggplot(tidy(Z), aes(index, value, color = series)) +
geom_line()
ggplot(tidy(Z), aes(index, value)) +
geom_line() +
facet_wrap(~series, ncol = 1)
Zrolled <- rollmean(Z, 5)
ggplot(tidy(Zrolled), aes(index, value, color = series)) +
geom_line()</pre>
```

tidy_irlba

Tidy a(n) irlba object masquerading as list

Description

Broom tidies a number of lists that are effectively S3 objects without a class attribute. For example, stats::optim(), svd() and interp::interp() produce consistent output, but because they do not have a class attribute, they cannot be handled by S3 dispatch.

These functions look at the elements of a list and determine if there is an appropriate tidying method to apply to the list. Those tidiers are implemented as functions of the form tidy_<function> or glance_<function> and are not exported (but they are documented!).

If no appropriate tidying method is found, they throw an error.

Usage

tidy_irlba(x, ...)

Arguments

| х | A list returned from irlba::irlba(). |
|---|--|
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | • tidy() methods will warn when supplied an exponentiate argument if it will be ignored. |
| | • augment() methods will warn when supplied a newdata argument if it will be ignored. |
| | |

Details

A very thin wrapper around tidy_svd().

tidy_irlba

Value

A tibble::tibble with columns depending on the component of PCA being tidied.

If matrix is "u", "samples", "scores", or "x" each row in the tidied output corresponds to the original data in PCA space. The columns are:

| row | ID of the original observation (i.e. rowname from original data). |
|-------|--|
| PC | Integer indicating a principal component. |
| value | The score of the observation for that particular principal component. That is, the location of the observation in PCA space. |

If matrix is "v", "rotation", "loadings" or "variables", each row in the tidied output corresponds to information about the principle components in the original space. The columns are:

| row | The variable labels (colnames) of the data set on which PCA was performed. | |
|---|--|--|
| PC | An integer vector indicating the principal component. | |
| value | The value of the eigenvector (axis score) on the indicated principal component. | |
| If matrix is "d", "eigenvalues" or "pcs", the columns are: | | |
| PC | An integer vector indicating the principal component. | |
| std.dev | Standard deviation explained by this PC. | |
| percent | Fraction of variation explained by this component (a numeric value between 0 and 1). | |
| cumulative Cumulative fraction of variation explained by principle components up component (a numeric value between 0 and 1). | | |

See Also

tidy(), irlba::irlba()

Other list tidiers: glance_optim(), list_tidiers, tidy_optim(), tidy_svd(), tidy_xyz() Other svd tidiers: augment.prcomp(), tidy.prcomp(), tidy_svd()

Examples

```
library(modeldata)
data(hpc_data)
mat <- scale(as.matrix(hpc_data[, 2:5]))
s <- svd(mat)
tidy_u <- tidy(s, matrix = "u")
tidy_u
tidy_d <- tidy(s, matrix = "d")
tidy_d</pre>
```

```
tidy_v <- tidy(s, matrix = "v")
tidy_v
library(ggplot2)
library(dplyr)
ggplot(tidy_d, aes(PC, percent)) +
  geom_point() +
  ylab("% of variance explained")
tidy_u %>%
  mutate(class = hpc_data$class[row]) %>%
  ggplot(aes(class, value)) +
  geom_boxplot() +
  facet_wrap(~PC, scale = "free_y")
```

tidy_optim

Tidy a(n) optim object masquerading as list

Description

Broom tidies a number of lists that are effectively S3 objects without a class attribute. For example, stats::optim(), svd() and interp::interp() produce consistent output, but because they do not have a class attribute, they cannot be handled by S3 dispatch.

These functions look at the elements of a list and determine if there is an appropriate tidying method to apply to the list. Those tidiers are implemented as functions of the form tidy_<function> or glance_<function> and are not exported (but they are documented!).

If no appropriate tidying method is found, they throw an error.

Usage

tidy_optim(x, ...)

Arguments

х

A list returned from stats::optim().

```
. . .
```

Additional arguments. Not used. Needed to match generic signature only. **Cautionary note:** Misspelled arguments will be absorbed in . . . , where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9, all computation will proceed using conf.level = 0.95. Two exceptions here are:

- tidy() methods will warn when supplied an exponentiate argument if it will be ignored.
- augment() methods will warn when supplied a newdata argument if it will be ignored.

tidy_svd

Value

A tibble::tibble() with columns:

| parameter | The parameter being modeled. | |
|-----------|---|--|
| std.error | The standard error of the regression term. | |
| value | The value/estimate of the component. Results from data reshaping. | |

std.error is only provided as a column if the Hessian is calculated.

Note

This function assumes that the provided objective function is a negative log-likelihood function. Results will be invalid if an incorrect function is supplied.

tidy(o) glance(o)

See Also

tidy(), stats::optim()

Other list tidiers: glance_optim(), list_tidiers, tidy_irlba(), tidy_svd(), tidy_xyz()

Examples

f <- function(x) (x[1] - 2)² + (x[2] - 3)² + (x[3] - 8)² o <- optim(c(1, 1, 1), f)</pre>

| tic | lv | svd |
|-----|-------|-----|
| | · J – | |

Tidy a(n) svd object masquerading as list

Description

Broom tidies a number of lists that are effectively S3 objects without a class attribute. For example, stats::optim(), svd() and interp::interp() produce consistent output, but because they do not have a class attribute, they cannot be handled by S3 dispatch.

These functions look at the elements of a list and determine if there is an appropriate tidying method to apply to the list. Those tidiers are implemented as functions of the form tidy_<function> or glance_<function> and are not exported (but they are documented!).

If no appropriate tidying method is found, they throw an error.

Usage

tidy_svd(x, matrix = "u", ...)

Arguments

| х | A list with components u, d, v returned by base::svd(). |
|--------|--|
| matrix | Character specifying which component of the PCA should be tidied. |
| | "u", "samples", "scores", or "x": returns information about the map from the original space into principle components space. "v", "rotation", "loadings" or "variables": returns information about the map from principle components space back into the original space. "d", "eigenvalues" or "pcs": returns information about the eigenvalues. |
| | Additional arguments. Not used. Needed to match generic signature only. Cautionary note: Misspelled arguments will be absorbed in, where they will be ignored. If the misspelled argument has a default value, the default value will be used. For example, if you pass conf.lvel = 0.9 , all computation will proceed using conf.level = 0.95 . Two exceptions here are: |
| | tidy() methods will warn when supplied an exponentiate argument if it will be ignored. augment() methods will warn when supplied a newdata argument if it will be ignored. |

Details

See https://stats.stackexchange.com/questions/134282/relationship-between-svd-and-pca-how-to-usesvd-to-perform-pca for information on how to interpret the various tidied matrices. Note that SVD is only equivalent to PCA on centered data.

Value

A tibble::tibble with columns depending on the component of PCA being tidied.

If matrix is "u", "samples", "scores", or "x" each row in the tidied output corresponds to the original data in PCA space. The columns are:

| row | ID of the original observation (i.e. rowname from original data). |
|-------|--|
| PC | Integer indicating a principal component. |
| value | The score of the observation for that particular principal component. That is, the location of the observation in PCA space. |

If matrix is "v", "rotation", "loadings" or "variables", each row in the tidied output corresponds to information about the principle components in the original space. The columns are:

| The variable labels (colnames) of the data set on which PCA was performed. | | | | |
|---|--|--|--|--|
| An integer vector indicating the principal component. | | | | |
| The value of the eigenvector (axis score) on the indicated principal component. | | | | |
| If matrix is "d", "eigenvalues" or "pcs", the columns are: | | | | |
| An integer vector indicating the principal component. | | | | |
| Standard deviation explained by this PC. | | | | |
| | | | | |

tidy_xyz

| percent | Fraction of variation explained by this component (a numeric value between 0 and 1). |
|------------|--|
| cumulative | Cumulative fraction of variation explained by principle components up to this component (a numeric value between 0 and 1). |

See Also

base::svd()

```
Other svd tidiers: augment.prcomp(), tidy.prcomp(), tidy_irlba()
Other list tidiers: glance_optim(), list_tidiers, tidy_irlba(), tidy_optim(), tidy_xyz()
```

Examples

```
library(modeldata)
data(hpc_data)
mat <- scale(as.matrix(hpc_data[, 2:5]))</pre>
s <- svd(mat)</pre>
tidy_u <- tidy(s, matrix = "u")</pre>
tidy_u
tidy_d <- tidy(s, matrix = "d")</pre>
tidy_d
tidy_v <- tidy(s, matrix = "v")</pre>
tidy_v
library(ggplot2)
library(dplyr)
ggplot(tidy_d, aes(PC, percent)) +
  geom_point() +
  ylab("% of variance explained")
tidy_u %>%
  mutate(class = hpc_data$class[row]) %>%
  ggplot(aes(class, value)) +
  geom_boxplot() +
  facet_wrap(~PC, scale = "free_y")
```

Description

Broom tidies a number of lists that are effectively S3 objects without a class attribute. For example, stats::optim(), svd() and interp::interp() produce consistent output, but because they do not have a class attribute, they cannot be handled by S3 dispatch.

These functions look at the elements of a list and determine if there is an appropriate tidying method to apply to the list. Those tidiers are implemented as functions of the form tidy_<function> or glance_<function> and are not exported (but they are documented!).

If no appropriate tidying method is found, they throw an error.

xyz lists (lists where x and y are vectors of coordinates and z is a matrix of values) are typically used by functions such as graphics::persp() or graphics::image() and returned by interpolation functions such as interp::interp().

Usage

tidy_xyz(x, ...)

Arguments

| Additional arguments. Not used. Needed to match generic signat tionary note: Misspelled arguments will be absorbed in, whi ignored. If the misspelled argument has a default value, the defau used. For example, if you pass conf.lvel = 0.9, all computation using conf.level = 0.95. Two exceptions here are: tidy() methods will warn when supplied an exponentiate will be ignored. augment() methods will warn when supplied a newdata arg be ignored. | nd z is a matrix. ength of y must |
|---|--|
| will be ignored.augment() methods will warn when supplied a newdata arg | nere they will be ult value will be |
| | e argument if it |
| be ignored. | gument if it will |

Value

A tibble::tibble with vector columns x, y and z.

See Also

tidy(), graphics::persp(), graphics::image(), interp::interp()
Other list tidiers: glance_optim(), list_tidiers, tidy_irlba(), tidy_optim(), tidy_svd()

Examples

```
A <- list(x = 1:5, y = 1:3, z = matrix(runif(5 * 3), nrow = 5))
image(A)
tidy(A)</pre>
```

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