Package 'gofreg'

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Title Bootstrap-Based Goodness-of-Fit Tests for Parametric Regression

Version 1.0.0

Description Provides statistical methods to check if a parametric family of conditional density functions fits to some given dataset of covariates and response variables. Different test statistics can be used to determine the goodness-of-fit of the assumed model, see Andrews (1997) <doi:10.2307/2171880>, Bierens & Wang (2012) <doi:10.1017/S0266466611000168>, Dikta & Scheer (2021) <doi:10.1007/978-3-030-73480-0> and Kremling & Dikta (2024) <doi:10.48550/arXiv.2409.20262>. As proposed in these papers, the corresponding p-values are approximated using a parametric bootstrap method.

```
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```

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Description

This class inherits from TestStatistic and implements a function to calculate the test statistic (and x-y-values that can be used to plot the underlying process).

The process underlying the test statistic is defined by

$$\bar{\alpha}_n(s,t) = \frac{1}{\sqrt{n}} \sum_{i=1}^n \left(I_{\{Y_i \le t\}} - F(t|\hat{\vartheta}_n, X_i) \right) I_{\{\hat{\beta}_n^T X_i \le s\}}, \quad (s,t) \in \mathbb{R}^2.$$

Super class

gofreg::TestStatistic -> CondKolmbXY

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Methods

Public methods:

• CondKolmbXY\$calc_stat()

deep Whether to make a deep clone.

• CondKolmbXY\$clone()

Method calc_stat(): Calculate the value of the test statistic for given data and a model to test for.

```
Usage:
CondKolmbXY$calc_stat(data, model)
Arguments:
data data.frame() with columns x and y containing the data
model ParamRegrModel to test for, already fitted to the data
Returns: The modified object (self), allowing for method chaining.

Method clone(): The objects of this class are cloneable with this method.
Usage:
CondKolmbXY$clone(deep = FALSE)
Arguments:
```

```
# Create an example dataset
n <- 100
x \leftarrow cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()</pre>
y <- model$sample_yx(x, params=list(beta=c(2,3), sd=1))</pre>
data \leftarrow dplyr::tibble(x = x, y = y)
# Fit the correct model
model$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Print value of test statistic and plot corresponding process
ts <- CondKolmbXY$new()</pre>
ts$calc_stat(data, model)
print(ts)
plot(ts)
# Fit a wrong model
model2 \leftarrow NormalGLM new(linkinv = function(u) {u+10})
model2$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Print value of test statistic and plot corresponding process
ts2 <- CondKolmbXY$new()
ts2$calc_stat(data, model2)
print(ts2)
plot(ts2)
```

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CondKolmXY

Conditional Kolmogorov test statistic for the joint distribution of (X,Y)

Description

This class inherits from TestStatistic and implements a function to calculate the test statistic (and x-y-values that can be used to plot the underlying process).

The process underlying the test statistic is given in Andrews (1997) doi: 10.2307/2171880 and defined by

$$\nu_n(x,y) = \frac{1}{\sqrt{n}} \sum_{i=1}^n \left(I_{\{Y_i \le y\}} - F(y|\hat{\vartheta}_n, X_i) \right) I_{\{X_i \le x\}}, \quad (x,y) \in \mathbb{R}^{p+1}.$$

Super class

gofreg::TestStatistic -> CondKolmXY

Methods

Public methods:

- CondKolmXY\$calc_stat()
- CondKolmXY\$clone()

Method calc_stat(): Calculate the value of the test statistic for given data and a model to test for

Usage:

CondKolmXY\$calc_stat(data, model)

Arguments:

data data.frame() with columns x and y containing the data model ParamRegrModel to test for, already fitted to the data

Returns: The modified object (self), allowing for method chaining.

Method clone(): The objects of this class are cloneable with this method.

Usage:

CondKolmXY\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

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Examples

```
# Create an example dataset
n <- 100
x \leftarrow cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()</pre>
y <- model$sample_yx(x, params=list(beta=c(2,3), sd=1))</pre>
data <- dplyr::tibble(x = x, y = y)
# Fit the correct model
model$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Print value of test statistic and plot corresponding process
ts <- CondKolmXY$new()</pre>
ts$calc_stat(data, model)
print(ts)
plot(ts)
# Fit a wrong model
model2 <- NormalGLM$new(linkinv = function(u) {u+10})</pre>
model2$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Print value of test statistic and plot corresponding process
ts2 <- CondKolmXY$new()
ts2$calc_stat(data, model2)
print(ts2)
plot(ts2)
```

CondKolmY

Conditional Kolmogorov test statistic for the marginal distribution of Y

Description

This class inherits from TestStatistic and implements a function to calculate the test statistic (and x-y-values that can be used to plot the underlying process).

The process underlying the test statistic is given in Kremling & Dikta (2024) https://arxiv.org/abs/2409.20262 and defined by

$$\tilde{\alpha}_n(t) = \frac{1}{\sqrt{n}} \sum_{i=1}^n \left(I_{\{Y_i \le t\}} - F(t|\hat{\vartheta}_n, X_i) \right), \quad -\infty \le t \le \infty.$$

Super class

```
gofreg::TestStatistic -> CondKolmY
```

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Methods

Public methods:

- CondKolmY\$calc_stat()
- CondKolmY\$clone()

Method calc_stat(): Calculate the value of the test statistic for given data and a model to test for.

```
Usage:
CondKolmY$calc_stat(data, model)
Arguments:
data data.frame() with columns x and y containing the data
model ParamRegrModel to test for, already fitted to the data
Returns: The modified object (self), allowing for method chaining.

Method clone(): The objects of this class are cloneable with this method.
Usage:
CondKolmY$clone(deep = FALSE)
Arguments:
deep Whether to make a deep clone.
```

```
# Create an example dataset
n <- 100
x \leftarrow cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()</pre>
y <- model$sample_yx(x, params=list(beta=c(2,3), sd=1))</pre>
data \leftarrow dplyr::tibble(x = x, y = y)
# Fit the correct model
model$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Print value of test statistic and plot corresponding process
ts <- CondKolmY$new()</pre>
ts$calc_stat(data, model)
print(ts)
plot(ts)
# Fit a wrong model
model2 \leftarrow NormalGLM new(linkinv = function(u) {u+10})
model2$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Print value of test statistic and plot corresponding process
ts2 <- CondKolmY$new()
ts2$calc_stat(data, model2)
print(ts2)
plot(ts2)
```

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CondKolmY_RCM

Conditional Kolmogorov test statistic for the marginal distribution of Y under random censorship

Description

This class inherits from TestStatistic and implements a function to calculate the test statistic (and x-y-values that can be used to plot the underlying process).

The process underlying the test statistic is defined by

$$\tilde{\alpha}_n^{KM}(t) = \sqrt{n} \left(\hat{F}_n^{KM}(t) - \frac{1}{n} \sum_{i=1}^n F(t | \hat{\vartheta}_n, X_i) \right), \quad -\infty \le t \le \infty.$$

Super class

```
gofreg::TestStatistic -> CondKolmY_RCM
```

Methods

Public methods:

- CondKolmY_RCM\$calc_stat()
- CondKolmY_RCM\$clone()

Method calc_stat(): Calculate the value of the test statistic for given data and a model to test for

Usage:

CondKolmY_RCM\$calc_stat(data, model)

Arguments:

data data.frame() with columns \boldsymbol{x} and \boldsymbol{y} containing the data

model ParamRegrModel to test for, already fitted to the data

Returns: The modified object (self), allowing for method chaining.

Method clone(): The objects of this class are cloneable with this method.

Usage.

CondKolmY_RCM\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

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Examples

```
# Create an example dataset
n <- 100
x <- cbind(runif(n), rbinom(n, 1, 0.5))</pre>
model <- NormalGLM$new()</pre>
y \leftarrow model\sample_yx(x, params=list(beta=c(2,3), sd=1))
c <- rnorm(n, mean(y)*1.2, sd(y)*0.5)
data <- dplyr::tibble(x = x, z = pmin(y,c), delta = as.numeric(y <= c))
# Fit the correct model
model$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE, loglik = loglik_xzd)
# Print value of test statistic and plot corresponding process
ts <- CondKolmY_RCM$new()</pre>
ts$calc_stat(data, model)
print(ts)
plot(ts)
# Fit a wrong model
model2 <- NormalGLM$new(linkinv = function(u) {u+10})</pre>
model2$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE, loglik = loglik_xzd)
# Print value of test statistic and plot corresponding process
ts2 <- CondKolmY_RCM$new()
ts2$calc_stat(data, model2)
print(ts2)
plot(ts2)
```

ExpGLM

Generalized linear model with exponential distribution

Description

This class represents a generalized linear model with exponential distribution. It inherits from GLM and implements its functions that, for example, evaluate the conditional density and distribution functions.

Super classes

```
gofreg::ParamRegrModel -> gofreg::GLM -> ExpGLM
```

Methods

Public methods:

- ExpGLM\$fit()
- ExpGLM\$f_yx()
- ExpGLM\$F_yx()
- ExpGLM\$F1_yx()

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```
• ExpGLM$sample_yx()
  • ExpGLM$clone()
Method fit(): Calculates the maximum likelihood estimator for the model parameters based
on given data.
 Usage:
 ExpGLM$fit(
    data,
    params_init = private$params,
   loglik = loglik_xy,
    inplace = FALSE
 Arguments:
 data tibble containing the data to fit the model to
 params_init initial value of the model parameters to use for the optimization (defaults to the
     fitted parameter values)
 loglik function(data, model, params) defaults to loglik_xy()
 inplace logical; if TRUE, default model parameters are set accordingly and parameter esti-
     mator is not returned
 Returns: MLE of the model parameters for the given data, same shape as params_init
Method f_yx(): Evaluates the conditional density function.
 Usage:
 ExpGLM$f_yx(t, x, params = private$params)
 Arguments:
 t value(s) at which the conditional density shall be evaluated
 x matrix of covariates, each row representing one sample
 params model parameters to use (list() with tag beta), defaults to the fitted parameter values
 Returns: value(s) of the conditional density function, same shape as t
Method F_yx(): Evaluates the conditional distribution function.
 ExpGLM$F_yx(t, x, params = private$params)
 Arguments:
 t value(s) at which the conditional distribution shall be evaluated
 x matrix of covariates, each row representing one sample
 params model parameters to use (list() with tag beta), defaults to the fitted parameter values
 Returns: value(s) of the conditional distribution function, same shape as t
Method F1_yx(): Evaluates the conditional quantile function.
 ExpGLM$F1_yx(t, x, params = private$params)
```

Arguments:

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t value(s) at which the conditional quantile function shall be evaluated x matrix of covariates, each row representing one sample params model parameters to use (list() with tag beta), defaults to the fitted parameter values *Returns:* value(s) of the conditional quantile function, same shape as t

Method sample_yx(): Generates a new sample of response variables with the same conditional distribution.

```
Usage:
ExpGLM$sample_yx(x, params = private$params)
Arguments:
x matrix of covariates, each row representing one sample
params model parameters to use (list() with tag beta), defaults to the fitted parameter values
Returns: vector of sampled response variables, same length as nrow(x)
```

Method clone(): The objects of this class are cloneable with this method.

```
Usage:
ExpGLM$clone(deep = FALSE)
Arguments:
deep Whether to make a deep clone.
```

```
# Use the built-in cars dataset
x <- datasets::cars$speed
y <- datasets::cars$dist</pre>
data <- dplyr::tibble(x=x, y=y)</pre>
# Create an instance of ExpGLM
model <- ExpGLM$new()</pre>
# Fit an Exponential GLM to the cars dataset
model$fit(data, params_init = list(beta=3), inplace=TRUE)
params_opt <- model$get_params()</pre>
# Plot the resulting regression function
plot(datasets::cars)
abline(a = 0, b = params_opt$beta)
# Generate a sample for y for given x following the same distribution
x.new \leftarrow seq(min(x), max(x), by=2)
y.smpl <- model$sample_yx(x.new)</pre>
points(x.new, y.smpl, col="red")
# Evaluate the conditional density, distribution, quantile and regression
# function at given values
model$f_yx(y.smpl, x.new)
model$F_yx(y.smpl, x.new)
model$F1_yx(y.smpl, x.new)
y.pred <- model$mean_yx(x.new)</pre>
points(x.new, y.pred, col="blue")
```

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GammaGLM

Generalized linear model with gamma distribution

Description

This class represents a generalized linear model with Gamma distribution. It inherits from GLM and implements its functions that, for example, evaluate the conditional density and distribution functions.

Super classes

```
gofreg::ParamRegrModel -> gofreg::GLM -> GammaGLM
```

Methods

Public methods:

- GammaGLM\$fit()
- GammaGLM\$f_yx()
- GammaGLM\$F_yx()
- GammaGLM\$F1_yx()
- GammaGLM\$sample_yx()
- GammaGLM\$clone()

Method fit(): Calculates the maximum likelihood estimator for the model parameters based on given data.

```
Usage:
GammaGLM$fit(
  data,
  params_init = private$params,
  loglik = loglik_xy,
  inplace = FALSE
)
Arguments:
```

data tibble containing the data to fit the model to

params_init initial value of the model parameters to use for the optimization (defaults to the fitted parameter values)

```
loglik function(data, model, params) defaults to loglik_xy()
```

inplace logical; if TRUE, default model parameters are set accordingly and parameter estimator is not returned

Returns: MLE of the model parameters for the given data, same shape as params_init

Method $f_yx()$: Evaluates the conditional density function.

```
Usage:
GammaGLM$f_yx(t, x, params = private$params)
```

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Arguments:

t value(s) at which the conditional density shall be evaluated

x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: value(s) of the conditional density function, same shape as t

Method F_yx(): Evaluates the conditional distribution function.

Usage:

```
GammaGLM$F_yx(t, x, params = private$params)
```

Arguments:

- t value(s) at which the conditional distribution shall be evaluated
- x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: value(s) of the conditional distribution function, same shape as t

Method F1_yx(): Evaluates the conditional quantile function.

Usage:

```
GammaGLM$F1_yx(t, x, params = private$params)
```

Arguments:

- t value(s) at which the conditional quantile function shall be evaluated
- x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: value(s) of the conditional quantile function, same shape as t

Method sample_yx(): Generates a new sample of response variables with the same conditional distribution.

Usage:

```
GammaGLM$sample_yx(x, params = private$params)
```

Arguments:

x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: vector of sampled response variables, same length as nrow(x)

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
GammaGLM$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

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Examples

```
# Use the built-in cars dataset
x <- datasets::cars$speed</pre>
v <- datasets::cars$dist</pre>
data <- dplyr::tibble(x=x, y=y)</pre>
# Create an instance of GammaGLM
model <- GammaGLM$new()</pre>
# Fit an Gamma GLM to the cars dataset
model$fit(data, params_init = list(beta=3, shape=1), inplace=TRUE)
params_opt <- model$get_params()</pre>
# Plot the resulting regression function
plot(datasets::cars)
abline(a = 0, b = params_opt$beta)
# Generate a sample for y for given x following the same distribution
x.new \leftarrow seq(min(x), max(x), by=2)
y.smpl <- model$sample_yx(x.new)</pre>
points(x.new, y.smpl, col="red")
# Evaluate the conditional density, distribution, quantile and regression
# function at given values
model$f_yx(y.smpl, x.new)
model$F_yx(y.smpl, x.new)
model$F1_yx(y.smpl, x.new)
y.pred <- model$mean_yx(x.new)</pre>
points(x.new, y.pred, col="blue")
```

GLM

Generalized linear model (abstract class)

Description

This class specializes ParamRegrModel. It is the abstract base class for parametric generalized linear model objects with specific distribution family such as NormalGLM and handles the (inverse) link function.

Super class

```
gofreg::ParamRegrModel -> GLM
```

Methods

Public methods:

- GLM\$new()
- GLM\$mean_yx()

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• GLM\$clone()

```
Method new(): Initialize an object of class GLM.
    Usage:
    GLM$new(linkinv = identity, params = NA)
    Arguments:
    linkinv inverse link function, defaults to identity function
```

params model parameters to use as default (optional)

Returns: a new instance of the class

Method mean_yx(): Evaluates the regression function or in other terms the expected value of Y given X=x.

```
Usage:
GLM$mean_yx(x, params = private$params)
Arguments:
x vector of covariates
params model parameters to use, defaults to the fitted parameter values
```

parameter values

Returns: value of the regression function

Method clone(): The objects of this class are cloneable with this method.

```
Usage:
GLM$clone(deep = FALSE)
Arguments:
```

deep Whether to make a deep clone.

GLM.new

Create GLM object with specific distribution family

Description

This constructor function can be used to create an instance of a parametric GLM with specific distribution family, returning a new object of NormalGLM, ExpGLM, WeibullGLM or GammaGLM, depending on the value of distr.

Usage

```
GLM.new(distr, linkinv = identity, params = NA)
```

Arguments

distr distribution family

linkinv inverse link function, defaults to identity function params model parameters to use as default (optional)

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Value

a new instance of a GLM-subclass

Examples

```
model <- GLM.new(distr = "normal")
# see examples of GLM-subclasses (e.g. NormalGLM) for how to use such models</pre>
```

GOFTest

Goodness-of-fit test for parametric regression

Description

This class implements functions to calculate the test statistic for the original data as well as the statistics for bootstrap samples. It also offers the possibility to compute the corresponding bootstrap p-value.

Methods

Public methods:

```
• GOFTest$new()
```

- GOFTest\$get_stat_orig()
- GOFTest\$get_stats_boot()
- GOFTest\$get_pvalue()
- GOFTest\$plot_procs()
- GOFTest\$clone()

Method new(): Initialize an instance of class GOFTest.

```
Usage:
GOFTest$new(
  data,
  model_fitted,
  test_stat,
  nboot,
  resample = resample_param,
  loglik = loglik_xy
)
Arguments:
data data.frame() containing the data
model_fitted object of class ParamRegrModel with fitted parameters
test_stat object of class TestStatistic
nboot number of bootstrap iterations
resample function(data, model) used to resample data in bootstrap iterations, defaults to
   resample_param()
```

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loglik function(data, model, params) negative log-likelihood function used to fit model

```
to resampled data in bootstrap iterations, defaults to loglik_xy()
 Returns: a new instance of the class
Method get_stat_orig(): Calculates the test statistic for the original data and model.
 Usage:
 GOFTest$get_stat_orig()
 Returns: object of class TestStatistic
Method get_stats_boot(): Calculates the test statistics for the resampled data and corre-
sponding models.
 Usage:
 GOFTest$get_stats_boot()
 Returns: vector of length nboot containing objects of class TestStatistic
Method get_pvalue(): Calculates the bootstrap p-value for the given model.
 Usage:
 GOFTest$get_pvalue()
 Returns: p-value for the null hypothesis that y was generated according to model
Method plot_procs(): Plots the processes underlying the bootstrap test statistics (gray) and
the original test statistic (red)
 Usage:
 GOFTest$plot_procs(
   title = sprintf("Test Statistic: %s, p-value: %s", class(private$test_stat)[1],
      self$get_pvalue()),
    subtitle = ggplot2::waiver(),
    color_boot = "gray40",
    color_orig = "red",
   x_{ab} = "plot.x",
    y_lab = "plot.y"
 )
 Arguments:
 title text to be displayed as title of the plot; defaults to "Test statistic: xxx, p-value: xxx"
 subtitle text to be displayed as subtitle of the plot; default is no subtitle
 color_boot color used to plot bootstrap test statistics; default is "red"
 color_orig color used to plot original test statistic; default is "gray40"
 x_lab label to use for the x-axis; default is "plot.x"
 y_lab label to use for the y-axis; default is "plot.y"
 Returns: The object (self), allowing for method chaining.
Method clone(): The objects of this class are cloneable with this method.
 Usage:
 GOFTest$clone(deep = FALSE)
 Arguments:
 deep Whether to make a deep clone.
```

loglik_xy

Examples

```
# Create an example dataset
n <- 100
x \leftarrow cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()</pre>
y <- model$sample_yx(x, params=list(beta=c(2,3), sd=1))</pre>
data <- dplyr::tibble(x = x, y = y)
# Fit the correct model
model$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Calculate the bootstrap p-value and plot the corresponding processes
goftest <- GOFTest$new(data, model, test_stat = CondKolmY$new(), nboot = 10)</pre>
goftest$get_pvalue()
goftest$plot_procs()
# Fit a wrong model
model2 <- NormalGLM$new(linkinv = function(u) {u+10})</pre>
model2$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Calculate the bootstrap p-value and plot the corresponding processes
goftest2 <- GOFTest$new(data, model2, test_stat = CondKolmY$new(), nboot = 10)</pre>
goftest2$get_pvalue()
goftest2$plot_procs()
```

loglik_xy

Negative log-likelihood function for a parametric regression model

Description

The log-likelihood function for a parametric regression model with data (x,y) is given by the sum of the logarithm of the conditional density of Y given X=x evaluated at y.

This function is one option that can be used to fit a ParamRegrModel. It returns the negative log-likelihood value in order for optim() to maximize (instead of minimize).

Usage

```
loglik_xy(data, model, params)
```

Arguments

data list() with tags x and y containing the data

model ParamRegrModel to use for the likelihood function

params vector with model parameters to compute likelihood function for

Value

Value of the negative log-likelihood function

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Examples

```
# Create an example dataset
n <- 100
x <- cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()
params.true <- list(beta = c(2,3), sd = 1)
y <- model$sample_yx(x, params = params.true)
data <- dplyr::tibble(x = x, y = y)

# Compute negative log likelihood for true parameters
loglik_xy(data, model, params.true)

# Compute negative log likelihood for wrong parameters (should be higher)
loglik_xy(data, model, params = list(beta = c(1,2), sd = 0.5))</pre>
```

loglik_xzd

Negative log-likelihood function for a parametric regression model under random censorship

Description

The log-likelihood function for a parametric regression model under random censorship with data (x,z,delta) is given by the sum of the logarithm of the conditional density of Y given X=x evaluated at z if z was uncensored or the logarithm of the conditional survival of Y given X=x evaluated at z if z was censored.

This function is one option that can be used to fit a ParamRegrModel. It returns the negative log-likelihood value in order for optim() to maximize (instead of minimize).

Usage

```
loglik_xzd(data, model, params)
```

Arguments

data list() with tags x, z and delta containing the data model ParamRegrModel to use for the likelihood function

params vector with model parameters to compute likelihood function for

Value

Value of the negative log-likelihood function

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Examples

```
# Create an example dataset
n <- 100
x <- cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()
params.true <- list(beta = c(2,3), sd = 1)
y <- model$sample_yx(x, params = params.true)
c <- rnorm(n, mean(y) * 1.2, sd(y) * 0.5)
data <- dplyr::tibble(x = x, z = pmin(y, c), delta = as.numeric(y <= c))
# Compute negative log likelihood for true parameters
loglik_xzd(data, model, params.true)
# Compute negative log likelihood for wrong parameters (should be higher)
loglik_xzd(data, model, params = list(beta = c(1,2), sd = 0.5))</pre>
```

MEP

Marked empirical process test statistic for a given GLM

Description

This class inherits from TestStatistic and implements a function to calculate the test statistic (and x-y-values that can be used to plot the underlying process).

The process underlying the test statistic is given in Dikta & Scheer (2021) doi: 10.1007/9783030-734800 and defined by

$$\bar{R}_n^1(u) = \frac{1}{\sqrt{n}} \sum_{i=1}^n \left(Y_i - m(X_i, \hat{\beta}_n) \right) I_{\{\hat{\beta}_n X_i \le u\}}, \quad -\infty \le u \le \infty.$$

Super class

```
gofreg::TestStatistic -> MEP
```

Methods

Public methods:

- MEP\$calc_stat()
- MEP\$clone()

Method calc_stat(): Calculate the value of the test statistic for given data and a model to test for.

```
Usage:
MEP$calc_stat(data, model)
Arguments:
data data.frame() with columns x and y containing the data
model ParamRegrModel to test for
```

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Returns: The modified object (self), allowing for method chaining.

Method clone(): The objects of this class are cloneable with this method.

```
Usage:
MEP$clone(deep = FALSE)
Arguments:
deep Whether to make a deep clone.
```

Examples

```
# Create an example dataset
n <- 100
x \leftarrow cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()</pre>
y <- model$sample_yx(x, params=list(beta=c(2,3), sd=1))</pre>
data \leftarrow dplyr::tibble(x = x, y = y)
# Fit the correct model
model$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Print value of test statistic and plot corresponding process
ts <- MEP$new()
ts$calc_stat(data, model)
print(ts)
plot(ts)
# Fit a wrong model
model2 <- NormalGLM$new(linkinv = function(u) {u+10})</pre>
model2$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Print value of test statistic and plot corresponding process
ts2 <- MEP$new()
ts2$calc_stat(data, model2)
print(ts2)
plot(ts2)
```

NegBinomGLM

Generalized linear model with negative binomial distribution

Description

This class represents a generalized linear model with negative binomial distribution. It inherits from GLM and implements its functions that, for example, evaluate the conditional density and distribution functions.

Super classes

```
gofreg::ParamRegrModel -> gofreg::GLM -> NegBinomGLM
```

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Methods

```
Public methods:
```

```
NegBinomGLM$fit()
NegBinomGLM$f_yx()
NegBinomGLM$F_yx()
NegBinomGLM$F1_yx()
NegBinomGLM$sample_yx()
NegBinomGLM$clone()
```

Method fit(): Calculates the maximum likelihood estimator for the model parameters based on given data.

```
Usage:
NegBinomGLM$fit(
    data,
    params_init = private$params,
    loglik = loglik_xy,
    inplace = FALSE
)
Arguments:
data tibble containing the data to fit the model to
params_init initial value of the model parameters to use for the optimization (defaults to the
    fitted parameter values)
loglik function(data, model, params) defaults to loglik_xy()
inplace logical; if TRUE, default model parameters are set accordingly and parameter esti-
mator is not returned
```

Returns: MLE of the model parameters for the given data, same shape as params_init

Method f_yx(): Evaluates the conditional density function.

```
Usage:
NegBinomGLM$f_yx(t, x, params = private$params)
Arguments:
t value(s) at which the conditional density shall be evaluated
```

x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: value(s) of the conditional density function, same shape as t

Method F_yx(): Evaluates the conditional distribution function.

```
Usage:
NegBinomGLM$F_yx(t, x, params = private$params)
Arguments:
t value(s) at which the conditional distribution shall be evaluated
x matrix of covariates, each row representing one sample
```

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params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: value(s) of the conditional distribution function, same shape as t

Method F1_yx(): Evaluates the conditional quantile function.

```
Usage:
```

```
NegBinomGLM$F1_yx(t, x, params = private$params)
```

Arguments:

- t value(s) at which the conditional quantile function shall be evaluated
- x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: value(s) of the conditional quantile function, same shape as t

Method sample_yx(): Generates a new sample of response variables with the same conditional distribution.

Usage:

```
NegBinomGLM$sample_yx(x, params = private$params)
```

Arguments.

x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: vector of sampled response variables, same length as nrow(x)

Method clone(): The objects of this class are cloneable with this method.

```
Usage:
```

```
NegBinomGLM$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

```
# Use the built-in cars dataset
x <- datasets::cars$speed
y <- datasets::cars$dist
data <- dplyr::tibble(x=x, y=y)

# Create an instance of a NegBinomGLM
model <- NegBinomGLM$new()

# Fit a Negative Binomial GLM to the cars dataset
model$fit(data, params_init = list(beta=3, shape=2), inplace=TRUE)
params_opt <- model$get_params()

# Plot the resulting regression function
plot(datasets::cars)</pre>
```

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```
abline(a = 0, b = params_opt$beta)

# Generate a sample for y for given x following the same distribution
x.new <- seq(min(x), max(x), by=2)
y.smpl <- model$sample_yx(x.new)
points(x.new, y.smpl, col="red")

# Evaluate the conditional density, distribution, quantile and regression
# function at given values
model$f_yx(y.smpl, x.new)
model$F_yx(y.smpl, x.new)
model$F1_yx(y.smpl, x.new)
y.pred <- model$mean_yx(x.new)
points(x.new, y.pred, col="blue")</pre>
```

NormalGLM

Generalized linear model with normal distribution

Description

This class represents a generalized linear model with normal distribution. It inherits from GLM and implements its functions that, for example, evaluate the conditional density and distribution functions.

Super classes

```
gofreg::ParamRegrModel -> gofreg::GLM -> NormalGLM
```

Methods

Public methods:

- NormalGLM\$fit()
- NormalGLM\$f_yx()
- NormalGLM\$F_yx()
- NormalGLM\$F1_yx()
- NormalGLM\$sample_yx()
- NormalGLM\$clone()

Method fit(): Calculates the maximum likelihood estimator for the model parameters based on given data.

```
Usage:
NormalGLM$fit(
  data,
  params_init = private$params,
  loglik = loglik_xy,
  inplace = FALSE
)
```

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Arguments:

data tibble containing the data to fit the model to

params_init initial value of the model parameters to use for the optimization (defaults to the fitted parameter values)

loglik function(data, model, params) defaults to loglik_xy()

inplace logical; if TRUE, default model parameters are set accordingly and parameter estimator is not returned

Returns: MLE of the model parameters for the given data, same shape as params_init

Method f_yx(): Evaluates the conditional density function.

Usage:

NormalGLM\$f_yx(t, x, params = private\$params)

Arguments:

- t value(s) at which the conditional density shall be evaluated
- x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and sd), defaults to the fitted parameter values

Returns: value(s) of the conditional density function, same shape as t

Method F_yx(): Evaluates the conditional distribution function.

Usage:

NormalGLM\$F_yx(t, x, params = private\$params)

Arguments:

- t value(s) at which the conditional distribution shall be evaluated
- x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and sd), defaults to the fitted parameter values

Returns: value(s) of the conditional distribution function, same shape as t

Method F1_yx(): Evaluates the conditional quantile function.

Usage:

NormalGLM\$F1_yx(t, x, params = private\$params)

Arguments:

- t value(s) at which the conditional quantile function shall be evaluated
- x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and sd), defaults to the fitted parameter values

Returns: value(s) of the conditional quantile function, same shape as t

Method sample_yx(): Generates a new sample of response variables with the same conditional distribution.

Usage:

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```
NormalGLM$sample_yx(x, params = private$params)

Arguments:

x matrix of covariates, each row representing one sample
params model parameters to use (list() with tags beta and sd), defaults to the fitted parameter
values

Returns: vector of sampled response variables, same length as nrow(x)

Method clone(): The objects of this class are cloneable with this method.

Usage:
```

Arguments:

deep Whether to make a deep clone.

NormalGLM\$clone(deep = FALSE)

```
# Use the built-in cars dataset
x <- datasets::cars$speed
y <- datasets::cars$dist</pre>
data <- dplyr::tibble(x=x, y=y)</pre>
# Create an instance of a NormalGLM
model <- NormalGLM$new()</pre>
# Fit a Normal GLM to the cars dataset
model$fit(data, params_init = list(beta=3, sd=2), inplace=TRUE)
params_opt <- model$get_params()</pre>
# Plot the resulting regression function
plot(datasets::cars)
abline(a = 0, b = params_opt$beta)
\# Generate a sample for y for given x following the same distribution
x.new \leftarrow seq(min(x), max(x), by=2)
y.smpl <- model$sample_yx(x.new)
points(x.new, y.smpl, col="red")
# Evaluate the conditional density, distribution, quantile and regression
# function at given values
model$f_yx(y.smpl, x.new)
model$F_yx(y.smpl, x.new)
model$F1_yx(y.smpl, x.new)
y.pred <- model$mean_yx(x.new)</pre>
points(x.new, y.pred, col="blue")
```

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ParamRegrModel

Parametric regression model (abstract class)

Description

This is the abstract base class for parametric regression model objects like NormalGLM.

Parametric regression models are built around the following key tasks:

- A method fit() to fit the model to given data, i.e. compute the MLE for the model parameters
- Methods f_yx(), F_yx() and mean_yx() to evaluate the conditional density, distribution and regression function
- A method sample_yx() to generate a random sample of response variables following the model given a vector of covariates

Methods

Public methods:

- ParamRegrModel\$set_params()
- ParamRegrModel\$get_params()
- ParamRegrModel\$fit()
- ParamRegrModel\$f_yx()
- ParamRegrModel\$F_yx()
- ParamRegrModel\$F1_yx()
- ParamRegrModel\$sample_yx()
- ParamRegrModel\$mean_yx()
- ParamRegrModel\$clone()

Method set_params(): Set the value of the model parameters used as default for the class functions.

Usage:

ParamRegrModel\$set_params(params)

Arguments:

params model parameters to use as default

Returns: The modified object (self), allowing for method chaining.

Method get_params(): Returns the value of the model parameters used as default for the class functions.

Usage:

ParamRegrModel\$get_params()

Returns: model parameters used as default

Method fit(): Calculates the maximum likelihood estimator for the model parameters based on given data.

```
Usage:
 ParamRegrModel$fit(data, params_init = private$params, loglik = loglik_xy)
 Arguments:
 data list containing the data to fit the model to
 params_init initial value of the model parameters to use for the optimization (defaults to the
     fitted parameter values)
 loglik function(data, model, params) defaults to loglik_xy()
 Returns: MLE of the model parameters for the given data, same shape as params_init
Method f_yx(): Evaluates the conditional density function.
 Usage:
 ParamRegrModel$f_yx(t, x, params = private$params)
 Arguments:
 t value(s) at which the conditional density shall be evaluated
 x vector of covariates
 params model parameters to use, defaults to the fitted parameter values
 Returns: value(s) of the conditional density function, same shape as t
Method F_yx(): Evaluates the conditional distribution function.
 Usage:
 ParamRegrModel$F_yx(t, x, params = private$params)
 Arguments:
 t value(s) at which the conditional distribution shall be evaluated
 x vector of covariates
 params model parameters to use, defaults to the fitted parameter values
 Returns: value(s) of the conditional distribution function, same shape as t
Method F1_yx(): Evaluates the conditional quantile function.
 Usage:
 ParamRegrModel$F1_yx(t, x, params = private$params)
 Arguments:
 t value(s) at which the conditional quantile function shall be evaluated
 x vector of covariates
 params model parameters to use, defaults to the fitted parameter values
 Returns: value(s) of the conditional quantile function, same shape as t
Method sample_yx(): Generates a new sample of response variables with the same conditional
distribution.
 Usage:
 ParamRegrModel$sample_yx(x, params = private$params)
 Arguments:
 x vector of covariates
```

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params model parameters to use, defaults to the fitted parameter values

Returns: vector of sampled response variables, same length as x

Method mean_yx(): Evaluates the regression function or in other terms the expected value of Y given X=x.

Usage:

ParamRegrModel\$mean_yx(x, params = private\$params)

Arguments:

x vector of covariates

params model parameters to use, defaults to the fitted parameter values

Returns: value of the regression function

Method clone(): The objects of this class are cloneable with this method.

Usage:

ParamRegrModel\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

resample_param

Parametric resampling scheme for a parametric regression model

Description

Generate a new, resampled dataset of the same shape as data following the given model. The covariates are kept the same and the response variables are drawn according to model\$sample_yx().

Usage

```
resample_param(data, model)
```

Arguments

data data.frame() with columns x and y containing the original data

model ParamRegrModel to use for the resampling

Value

data.frame() with columns x and y containing the resampled data

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Examples

```
# Create an example dataset
n <- 10
x <- cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()
params <- list(beta = c(2, 3), sd = 1)
y <- model$sample_yx(x, params = params)
data <- dplyr::tibble(x = x, y = y)

# Fit the model to the data
model$fit(data, params_init = params, inplace = TRUE)
# Resample from the model given data
resample_param(data, model)</pre>
```

resample_param_cens

Parametric resampling scheme for a parametric regression model under random censorship

Description

Generate a new, resampled dataset of the same shape as data following the given model. The covariates X are kept the same. Survival times Y are drawn according to model $sample_yx()$ and censoring times Y according to the Y are drawn according to model $sample_yx()$ and censoring times Y according to the Y are drawn according to modely and y are dra

Usage

```
resample_param_cens(data, model)
```

Arguments

data data.frame() with columns x, z and delta containing the original data

model ParamRegrModel to use for the resampling

Value

data.frame() with columns x, z and delta containing the resampled data

```
# Create an example dataset
n <- 10
x <- cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()
params <- list(beta = c(2, 3), sd = 1)
y <- model$sample_yx(x, params = params)
c <- rnorm(n, mean(y) * 1.2, sd(y) * 0.5)
z <- pmin(y, c)
delta <- as.numeric(y <= c)</pre>
```

```
data <- dplyr::tibble(x = x, z = z, delta = delta)
# Fit the model to the data
model$fit(data, params_init = params, inplace = TRUE, loglik = loglik_xzd)
# Resample from the model given data
resample_param_cens(data, model)</pre>
```

Description

Generate a new, resampled dataset of the same shape as data following the given model. The covariates are resampled from data\$x and the response variables are drawn according to model\$sample_yx().

Usage

```
resample_param_rsmplx(data, model)
```

Arguments

data data frame() with columns x and y containing the original data

model ParamRegrModel to use for the resampling

Value

data.frame() with columns x and y containing the resampled data

```
# Create an example dataset
n <- 10
x <- cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()
params <- list(beta = c(2, 3), sd = 1)
y <- model$sample_yx(x, params = params)
data <- dplyr::tibble(x = x, y = y)

# Fit the model to the data
model$fit(data, params_init = params, inplace = TRUE)
# Resample from the model given data
resample_param(data, model)</pre>
```

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SICM

Simulated integrated conditional moment test statistic

Description

This class inherits from TestStatistic and implements a function to calculate the test statistic (and x-y-values that can be used to plot the underlying process).

The process underlying the test statistic is given in Bierens & Wang (2012) doi: 10.1017/S0266466611000168 and defined by

$$\hat{T}_{n}^{(s)}(c) = \frac{1}{(2c)^{p+1}} \int_{[-c,c]^{p}}^{c} \int_{-c}^{c} \left| \frac{1}{\sqrt{n}} \sum_{j=1}^{n} \left(\exp(i\tau Y_{j}) - \exp(i\tau \tilde{Y}_{j}) \right) \exp(i\xi^{T} X_{j}) \right|^{2} d\tau d\xi$$

Super class

```
gofreg::TestStatistic -> SICM
```

Methods

Public methods:

• SICM\$new()

Arguments:

- SICM\$calc_stat()
- SICM\$clone()

Method new(): Initialize an instance of class SICM.

```
Usage:
SICM$new(
  С,
  transx = function(values) {
     tvals <- atan(scale(values))</pre>
     tvals[,
    apply(values, 2, sd) == 0] <- 0
     return(tvals)
},
  transy = function(values, data) {
     array(atan(scale(values, center = mean(data$y),
    scale = sd(data$y))))
}
)
```

c chosen value for integral boundaries (see Bierens & Wang (2012))

transx function(values) used to transform x-values to be standardized and bounded; default is standardization by subtracting the mean and dividing by the standard deviation and then applying arctan

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transy function(values, data) used to transform y-values to be standardized and bounded (same method is used for simulated y-values); default is standardization by subtracting the mean and dividing by the standard deviation and then applying arctan

Returns: a new instance of the class

Method calc_stat(): Calculate the value of the test statistic for given data and a model to test for.

```
Usage:
SICM$calc_stat(data, model)
Arguments:
data data.frame() with columns x and y containing the data
model ParamRegrModel to test for
Returns: The modified object (self), allowing for method chaining.

Method clone(): The objects of this class are cloneable with this method.
Usage:
SICM$clone(deep = FALSE)
Arguments:
deep Whether to make a deep clone.
```

```
# Create an example dataset
n <- 100
x \leftarrow cbind(runif(n), rbinom(n, 1, 0.5))
model <- NormalGLM$new()</pre>
y <- model$sample_yx(x, params=list(beta=c(2,3), sd=1))</pre>
data \leftarrow dplyr::tibble(x = x, y = y)
# Fit the correct model
model$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Print value of test statistic and plot corresponding process
ts <- SICM$new(c = 5)
ts$calc_stat(data, model)
print(ts)
plot(ts)
# Fit a wrong model
model2 <- NormalGLM$new(linkinv = function(u) {u+10})</pre>
model2$fit(data, params_init=list(beta=c(1,1), sd=3), inplace = TRUE)
# Print value of test statistic and plot corresponding process
ts2 <- SICM$new(c = 5)
ts2$calc_stat(data, model2)
print(ts2)
plot(ts2)
```

TestStatistic 33

TestStatistic

Test Statistic for parametric regression models (abstract class)

Description

This is the abstract base class for test statistic objects like CondKolmY or MEP.

Test statistics are built around the key method calc_stat() which calculates the particular test statistic (and x-y-values that can be used to plot the underlying process).

Methods

Public methods:

- TestStatistic\$get_value()
- TestStatistic\$calc_stat()
- TestStatistic\$get_plot_xy()
- TestStatistic\$print()
- TestStatistic\$geom_ts_proc()
- TestStatistic\$plot()
- TestStatistic\$clone()

Method get_value(): Returns the value of the test statistic.

Usage:

TestStatistic\$get_value()

Returns: value of the test statistic

Method calc_stat(): Calculate the value of the test statistic for given data and a model to test for.

Usage:

TestStatistic\$calc_stat(data, model)

Arguments:

data list() containing the data

model ParamRegrModel to test for

Returns: The modified object (self), allowing for method chaining.

Method $get_plot_xy()$: Returns vectors of x and y that can be used to plot the process corresponding to the test statistic.

Usage:

TestStatistic\$get_plot_xy()

Returns: list with plot.x and plot.y being vectors of the same length

Method print(): Overrides the print-method for objects of type TestStatistic to only print its value.

```
Usage:
 TestStatistic$print()
 Returns: The object (self), allowing for method chaining.
Method geom_ts_proc(): Creates a line plot showing the underlying process of the test statistic.
 Usage:
 TestStatistic$geom_ts_proc(...)
 Arguments:
 ... Other arguments passed on to ggplot2::geom_line(). These are often aesthetics, used
     to set an aesthetic to a fixed value, like colour = "red" or size = 3.
 Returns: A ggplot2 layer representing a line plot.
Method plot(): Creates a new ggplot showing the underlying process of the test statistic.
 Usage:
 TestStatistic$plot(...)
 Arguments:
 ... Other arguments passed on to ggplot2::geom_line(). These are often aesthetics, used
     to set an aesthetic to a fixed value, like colour = "red" or size = 3.
 Returns: A ggplot2 object representing the complete plot, including a line geometry.
Method clone(): The objects of this class are cloneable with this method.
 Usage:
 TestStatistic$clone(deep = FALSE)
 Arguments:
```

WeibullGLM

Generalized linear model with Weibull distribution

Description

This class represents a generalized linear model with Weibull distribution. It inherits from GLM and implements its functions that, for example, evaluate the conditional density and distribution functions.

Super classes

```
gofreg::ParamRegrModel -> gofreg::GLM -> WeibullGLM
```

deep Whether to make a deep clone.

Methods

```
Public methods:
```

```
WeibullGLM$fit()
WeibullGLM$f_yx()
WeibullGLM$F_yx()
WeibullGLM$F1_yx()
WeibullGLM$sample_yx()
WeibullGLM$clone()
```

Method fit(): Calculates the maximum likelihood estimator for the model parameters based on given data.

```
Usage:
WeibullGLM$fit(
    data,
    params_init = private$params,
    loglik = loglik_xy,
    inplace = FALSE
)

Arguments:
data tibble containing the data to fit the model to
    params_init initial value of the model parameters to use for the optimization (defaults to the
        fitted parameter values)
loglik function(data, model, params) defaults to loglik_xy()
inplace logical; if TRUE, default model parameters are set accordingly and parameter esti-
        mator is not returned
```

Returns: MLE of the model parameters for the given data, same shape as params_init

Method $f_yx()$: Evaluates the conditional density function.

```
Usage:
WeibullGLM$f_yx(t, x, params = private$params)
Arguments:
t value(s) at which the conditional density shall be evaluated
x matrix of covariates, each row representing one sample
params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values
```

Returns: value(s) of the conditional density function, same shape as t

Method F_yx(): Evaluates the conditional distribution function.

```
Usage:
WeibullGLM$F_yx(t, x, params = private$params)
Arguments:
t value(s) at which the conditional distribution shall be evaluated
x matrix of covariates, each row representing one sample
```

params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: value(s) of the conditional distribution function, same shape as t

Method F1_yx(): Evaluates the conditional quantile function.

Usage:

```
WeibullGLM$F1_yx(t, x, params = private$params)
```

Arguments:

- t value(s) at which the conditional quantile function shall be evaluated
- x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: value(s) of the conditional quantile function, same shape as t

Method sample_yx(): Generates a new sample of response variables with the same conditional distribution.

Usage:

```
WeibullGLM$sample_yx(x, params = private$params)
```

Arguments

x matrix of covariates, each row representing one sample

params model parameters to use (list() with tags beta and shape), defaults to the fitted parameter values

Returns: vector of sampled response variables, same length as nrow(x)

Method clone(): The objects of this class are cloneable with this method.

Usage

```
WeibullGLM$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

```
# Use the built-in cars dataset
x <- datasets::cars$speed
y <- datasets::cars$dist
data <- dplyr::tibble(x=x, y=y)

# Create an instance of WeibullGLM
model <- WeibullGLM$new()

# Fit an Weibull GLM to the cars dataset
model$fit(data, params_init = list(beta=3, shape=1), inplace=TRUE)
params_opt <- model$get_params()

# Plot the resulting regression function
plot(datasets::cars)</pre>
```

```
abline(a = 0, b = params_opt$beta)

# Generate a sample for y for given x following the same distribution
x.new <- seq(min(x), max(x), by=2)
y.smpl <- model$sample_yx(x.new)
points(x.new, y.smpl, col="red")

# Evaluate the conditional density, distribution, quantile and regression
# function at given values
model$f_yx(y.smpl, x.new)
model$F_yx(y.smpl, x.new)
model$F1_yx(y.smpl, x.new)
y.pred <- model$mean_yx(x.new)
points(x.new, y.pred, col="blue")</pre>
```

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