

Package ‘fastcpd’

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Type Package

Title Fast Change Point Detection via Sequential Gradient Descent

Version 0.16.2

Description Implements fast change point detection algorithm based on the paper ``Sequential Gradient Descent and Quasi-Newton's Method for Change-Point Analysis'' by Xianyang Zhang, Trisha Dawn <<https://proceedings.mlr.press/v206/zhang23b.html>>. The algorithm is based on dynamic programming with pruning and sequential gradient descent. It is able to detect change points a magnitude faster than the vanilla Pruned Exact Linear Time(PELT). The package includes examples of linear regression, logistic regression, Poisson regression, penalized linear regression data, and whole lot more examples with custom cost function in case the user wants to use their own cost function.

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URL <https://fastcpd.xingchi.li>, <https://github.com/doccstat/fastcpd>

BugReports <https://github.com/doccstat/fastcpd/issues>

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Contents

bitcoin	3
fastcpd	4
fastcpd-class	11
fastcpd_ar	12
fastcpd_arima	13
fastcpd_arma	14
fastcpd_binomial	16
fastcpd_garch	17
fastcpd_lasso	18
fastcpd_lm	20
fastcpd_mean	21
fastcpd_meanvariance	22
fastcpd_poisson	23
fastcpd_ts	24
fastcpd_var	27
fastcpd_variance	28
occupancy	29
plot.fastcpd	30
print.fastcpd	32
show.fastcpd	33
summary.fastcpd	33
transcriptome	34
uk_seatbelts	36
variance_arma	38
variance_lm	39
variance_mean	40
variance_median	41
well_log	41

bitcoin	<i>Bitcoin Market Price (USD)</i>
---------	-----------------------------------

Description

The average USD market price across major bitcoin exchanges.

Usage

```
bitcoin
```

Format

A data frame with 1354 rows and 2 variables:

date POSIXct,POSIXt (TZ: "UTC") from 2019-01-02 to 2023-10-28
price The average USD market price across major bitcoin exchanges

Source

<<https://www.blockchain.com/explorer/charts/market-price>>

Examples

```
if (requireNamespace("ggplot2", quietly = TRUE)) {  
  p <- ggplot2::ggplot(bitcoin, ggplot2::aes(x = date, y = price)) +  
    ggplot2::geom_line()  
  print(p)  
  
  result <- suppressWarnings(fastcpd.garch(  
    diff(log(bitcoin$price[600:900])), c(1, 1),  
    beta = "BIC", cost_adjustment = "BIC"  
  ))  
  summary(result)  
  bitcoin$date[result@cp_set + 600]  
  plot(result)  
  
  cp_dates <- bitcoin[600 + result@cp_set + 1, "date"]  
  ggplot2::ggplot(  
    data = data.frame(  
      x = bitcoin$date[600:900], y = bitcoin$price[600:900]  
    ),  
    ggplot2::aes(x = x, y = y)  
  ) +  
    ggplot2::geom_line(color = "steelblue") +  
    ggplot2::geom_vline(  
      xintercept = cp_dates,  
      color = "red",  
      linetype = "dotted",
```

```

    linewidth = 0.5,
    alpha = 0.7
) +
ggplot2::labs(
  x = "Year",
  y = "Bitcoin price in USD"
) +
ggplot2::annotate(
  "text",
  x = cp_dates,
  y = 2000,
  label = as.character(cp_dates),
  color = "steelblue"
) +
ggplot2::theme_bw()
}

```

fastcpd*Find change points efficiently***Description**

fastcpd() takes in formulas, data, families and extra parameters and returns a **fastcpd** object.

Usage

```

fastcpd(
  formula = y ~ . ~ 1,
  data,
  beta = "MBIC",
  cost_adjustment = "MBIC",
  family = NULL,
  cost = NULL,
  cost_gradient = NULL,
  cost_hessian = NULL,
  line_search = c(1),
  lower = rep(-Inf, p),
  upper = rep(Inf, p),
  pruning_coef = 0,
  segment_count = 10,
  trim = 0.05,
  momentum_coef = 0,
  multiple_epochs = function(x) 0,
  epsilon = 1e-10,
  order = c(0, 0, 0),
  p = ncol(data) - 1,
  variance_estimation = NULL,

```

```

cp_only = FALSE,
vanilla_percentage = 0,
warm_start = FALSE,
...
)

```

Arguments

<code>formula</code>	A formula object specifying the model to be fitted. The (optional) response variable should be on the LHS of the formula, while the covariates should be on the RHS. The naming of variables used in the formula should be consistent with the column names in the data frame provided in <code>data</code> . The intercept term should be removed from the formula. The response variable is not needed for mean/variance change models and time series models. By default, an intercept column will be added to the data, similar to the <code>lm()</code> function. Thus, it is suggested that users should remove the intercept term by appending <code>- 1</code> to the formula. Note that the <code>fastcpd.family</code> functions do not require a formula input.
<code>data</code>	A data frame of dimension $T \times d$ containing the data to be segmented (where each row denotes a data point $z_t \in \mathbb{R}^d$ for $t = 1, \dots, T$) is required in the main function, while a matrix or a vector input is also accepted in the <code>fastcpd.family</code> functions.
<code>beta</code>	Penalty criterion for the number of change points. This parameter takes a string value of "BIC", "MBIC", "MDL" or a numeric value. If a numeric value is provided, the value will be used as the penalty. By default, the mBIC criterion is used, where $\beta = (p + 2) \log(T)/2$. This parameter usage should be paired with <code>cost_adjustment</code> described below. Discussions about the penalty criterion can be found in the references.
<code>cost_adjustment</code>	Cost adjustment criterion. It can be "BIC", "MBIC", "MDL" or NULL. By default, the cost adjustment criterion is set to be "MBIC". The "MBIC" and "MDL" criteria modify the cost function by adding a negative adjustment term to the cost function. "BIC" or NULL does not modify the cost function. Details can be found in the references.
<code>family</code>	Family class of the change point model. It can be "mean" for mean change, "variance" for variance change, "meanvariance" for mean and/or variance change, "lm" for linear regression, "binomial" for logistic regression, "poisson" for Poisson regression, "lasso" for penalized linear regression, "ar" for AR(p) models, "arma" for ARMA(p, q) models, "arima" for ARIMA(p, d, q) models, "garch" for GARCH(p, q) models, "var" for VAR(p) models and "custom" for user-specified custom models. Omitting this parameter is the same as specifying the parameter to be "custom" or NULL, in which case, users must specify the custom cost function.
<code>cost</code>	Cost function to be used. <code>cost</code> , <code>cost_gradient</code> , and <code>cost_hessian</code> should not be specified at the same time with <code>family</code> as built-in families have cost functions implemented in C++ to provide better performance. If not specified, the default is the negative log-likelihood for the corresponding family. Custom cost functions can be provided in the following two formats:

- `cost = function(data) { ... }`
- `cost = function(data, theta) { ... }`

Users can specify a loss function using the second format that will be used to calculate the cost value. In both formats, the input data is a subset of the original data frame in the form of a matrix (a matrix with a single column in the case of a univariate data set). In the first format, the specified cost function directly calculates the cost value. `fastcpd()` performs the vanilla PELT algorithm, and `cost_gradient` and `cost_hessian` should not be provided since no parameter updating is necessary for vanilla PELT. In the second format, the loss function $\sum_{i=s}^t l(z_i, \theta)$ is provided, which has to be optimized over the parameter θ to obtain the cost value. A detailed discussion about the custom cost function usage can be found in the references.

`cost_gradient` Gradient of the custom cost function. Example usage:

```
cost_gradient = function(data, theta) {
  ...
  return(gradient)
}
```

The gradient function takes two inputs, the first being a matrix representing a segment of the data, similar to the format used in the `cost` function, and the second being the parameter that needs to be optimized. The gradient function returns the value of the gradient of the loss function, i.e., $\sum_{i=s}^t \nabla l(z_i, \theta)$.

`cost_hessian` Hessian of the custom loss function. The Hessian function takes two inputs, the first being a matrix representing a segment of the data, similar to the format used in the `cost` function, and the second being the parameter that needs to be optimized. The gradient function returns the Hessian of the loss function, i.e., $\sum_{i=s}^t \nabla^2 l(z_i, \theta)$.

`line_search` If a vector of numeric values is provided, a line search will be performed to find the optimal step size for each update. Detailed usage of `line_search` can be found in the references.

`lower` Lower bound for the parameters. Used to specify the domain of the parameters after each gradient descent step. If not specified, the lower bound is set to be `-Inf` for all parameters. `lower` is especially useful when the estimated parameters take only positive values, such as the noise variance.

`upper` Upper bound for the parameters. Used to specify the domain of the parameters after each gradient descent step. If not specified, the upper bound is set to be `Inf` for all parameters.

`pruning_coeff` Pruning coefficient `c_0$` used in the pruning step of the PELT algorithm with the default value 0. If `cost_adjustment` is specified as "MBIC", an adjustment term $p \log(2)$ will be added to the pruning coefficient. If `cost_adjustment` is specified as "MDL", an adjustment term $p \log_2(2)$ will be added to the pruning coefficient. Detailed discussion about the pruning coefficient can be found in the references.

`segment_count` An initial guess of the number of segments. If not specified, the initial guess of the number of segments is 10. The initial guess affects the initial estimates of the parameters in SeGD.

<code>trim</code>	Trimming for the boundary change points so that a change point close to the boundary will not be counted as a change point. This parameter also specifies the minimum distance between two change points. If several change points have mutual distances smaller than <code>trim * nrow(data)</code> , those change points will be merged into one single change point. The value of this parameter should be between 0 and 1.
<code>momentum_coef</code>	Momentum coefficient to be applied to each update. This parameter is used when the loss function is bad-shaped so that maintaining a momentum from previous update is desired. Default value is 0, meaning the algorithm doesn't maintain a momentum by default.
<code>multiple_epochs</code>	A function can be specified such that an adaptive number of multiple epochs can be utilized to improve the algorithm's performance. <code>multiple_epochs</code> is a function of the length of the data segment. The function returns an integer indicating how many epochs should be performed apart from the default update. By default, the function returns zero, meaning no multiple epochs will be used to update the parameters. Example usage:
	<pre>multiple_epochs = function(segment_length) { if (segment_length < 100) 1 else 0 }</pre>
	This function will let SeGD perform parameter updates with an additional epoch for each segment with a length less than 100 and no additional epoch for segments with lengths greater or equal to 100.
<code>epsilon</code>	Epsilon to avoid numerical issues. Only used for the Hessian computation in Logistic Regression and Poisson Regression.
<code>order</code>	Order of the AR(p), VAR(p) or ARIMA(p, d, q) model.
<code>p</code>	Number of covariates in the model. If not specified, the number of covariates will be inferred from the data, i.e., $p = \text{ncol}(\text{data}) - 1$. This parameter is superseded by <code>order</code> in the case of time series models: "ar", "var", "arima".
<code>variance_estimation</code>	An estimate of the variance / covariance matrix for the data. If not specified, the variance / covariance matrix will be estimated using the data.
<code>cp_only</code>	If TRUE, only the change points are returned. Otherwise, the cost function values together with the estimated parameters for each segment are also returned. By default the value is set to be FALSE so that <code>plot</code> can be used to visualize the results for a built-in model. <code>cp_only</code> has some performance impact on the algorithm, since the cost values and estimated parameters for each segment need to be calculated and stored. If the users are only interested in the change points, setting <code>cp_only</code> to be TRUE will help with the computational cost.
<code>vanilla_percentage</code>	The parameter v is between zero and one. For each segment, when its length is no more than vT , the cost value will be computed by performing an exact minimization of the loss function over the parameter. When its length is greater than vT , the cost value is approximated through SeGD. Therefore, this parameter induces an algorithm that can be interpreted as an interpolation between dynamic

programming with SeGD ($v = 0$) and the vanilla PELT ($v = 1$). The readers are referred to the references for more details.

warm_start

If TRUE, the algorithm will use the estimated parameters from the previous segment as the initial value for the current segment. This parameter is only used for the "glm" families.

...

Other parameters for specific models.

- `include.mean` is used to determine if a mean intercept term should be included in the ARIMA(p, d, q) or GARCH(p, q) models.
- `r.progress` is used to control the progress bar. By default the progress bar will be shown. To disable it, set `r.progress = FALSE`.
- `p.response` is used to specify the number of response variables. This parameter is especially useful for linear models with multivariate responses.

Value

A `fastcpd` object.

Gallery

<https://github.com/doccstat/fastcpd/tree/main/tests/testthat/examples>

References

Xingchi Li, Xianyang Zhang (2024). “fastcpd: Fast Change Point Detection in R.” *arXiv:2404.05933*, <https://arxiv.org/abs/2404.05933>.

Xianyang Zhang, Trisha Dawn (2023). “Sequential Gradient Descent and Quasi-Newton’s Method for Change-Point Analysis.” In Ruiz, Francisco, Dy, Jennifer, van de Meent, Jan-Willem (eds.), *Proceedings of The 26th International Conference on Artificial Intelligence and Statistics*, volume 206 series Proceedings of Machine Learning Research, 1129-1143.

See Also

`fastcpd.family` for the family-specific function; `plot.fastcpd()` for plotting the results, `summary.fastcpd()` for summarizing the results.

Examples

```
if (requireNamespace("mvtnorm", quietly = TRUE)) {
  set.seed(1)
  n <- 200
  p <- 4
  d <- 2
  x <- mvtnorm::rmvnorm(n, rep(0, p), diag(p))
  theta_1 <- matrix(runif(8, -3, -1), nrow = p)
  theta_2 <- matrix(runif(8, -1, 3), nrow = p)
  y <- rbind(
    x[1:125, ] %*% theta_1 + mvtnorm::rmvnorm(125, rep(0, d), 3 * diag(d)),
    x[126:n, ] %*% theta_2 + mvtnorm::rmvnorm(75, rep(0, d), 3 * diag(d))
  )
```

```

result_mlm <- fastcpd(
  cbind(y.1, y.2) ~ . - 1, cbind.data.frame(y = y, x = x), family = "lm"
)
summary(result_mlm)
}
if (
  requireNamespace("mvtnorm", quietly = TRUE) &&
  requireNamespace("stats", quietly = TRUE)
) {
  set.seed(1)
  n <- 400 + 300 + 500
  p <- 5
  x <- mvtnorm::rmvnorm(n, mean = rep(0, p), sigma = diag(p))
  theta <- rbind(
    mvtnorm::rmvnorm(1, mean = rep(0, p - 3), sigma = diag(p - 3)),
    mvtnorm::rmvnorm(1, mean = rep(5, p - 3), sigma = diag(p - 3)),
    mvtnorm::rmvnorm(1, mean = rep(9, p - 3), sigma = diag(p - 3))
  )
  theta <- cbind(theta, matrix(0, 3, 3))
  theta <- theta[rep(seq_len(3), c(400, 300, 500)), ]
  y_true <- rowSums(x * theta)
  factor <- c(
    2 * stats::rbinom(400, size = 1, prob = 0.95) - 1,
    2 * stats::rbinom(300, size = 1, prob = 0.95) - 1,
    2 * stats::rbinom(500, size = 1, prob = 0.95) - 1
  )
  y <- factor * y_true + stats::rnorm(n)
  data <- cbind.data.frame(y, x)
  huber_threshold <- 1
  huber_loss <- function(data, theta) {
    residual <- data[, 1] - data[, -1, drop = FALSE] %*% theta
    indicator <- abs(residual) <= huber_threshold
    sum(
      residual^2 / 2 * indicator +
      huber_threshold * (
        abs(residual) - huber_threshold / 2
      ) * (1 - indicator)
    )
  }
  huber_loss_gradient <- function(data, theta) {
    residual <- c(data[nrow(data), 1] - data[nrow(data), -1] %*% theta)
    if (abs(residual) <= huber_threshold) {
      -residual * data[nrow(data), -1]
    } else {
      -huber_threshold * sign(residual) * data[nrow(data), -1]
    }
  }
  huber_loss_hessian <- function(data, theta) {
    residual <- c(data[nrow(data), 1] - data[nrow(data), -1] %*% theta)
    if (abs(residual) <= huber_threshold) {
      outer(data[nrow(data), -1], data[nrow(data), -1])
    } else {
      0.01 * diag(length(theta))
    }
  }
}

```

```

        }
    }
huber_regression_result <- fastcpd(
  formula = y ~ . - 1,
  data = data,
  beta = (p + 1) * log(n) / 2,
  cost = huber_loss,
  cost_gradient = huber_loss_gradient,
  cost_hessian = huber_loss_hessian
)
summary(huber_regression_result)
}

set.seed(1)
p <- 1
x <- matrix(rnorm(375 * p, 0, 1), ncol = p)
theta <- rbind(rnorm(p, 0, 1), rnorm(p, 2, 1))
y <- c(
  rbinom(200, 1, 1 / (1 + exp(-x[1:200, ] %*% theta[1, , drop = FALSE]))),
  rbinom(175, 1, 1 / (1 + exp(-x[201:375, ] %*% theta[2, , drop = FALSE])))
)
data <- data.frame(y = y, x = x)
result_builtin <- suppressWarnings(fastcpd.binomial(data))
logistic_loss <- function(data, theta) {
  x <- data[, -1, drop = FALSE]
  y <- data[, 1]
  u <- x %*% theta
  nll <- -y * u + log(1 + exp(u))
  nll[u > 10] <- -y[u > 10] * u[u > 10] + u[u > 10]
  sum(nll)
}
logistic_loss_gradient <- function(data, theta) {
  x <- data[nrow(data), -1, drop = FALSE]
  y <- data[nrow(data), 1]
  c(-(y - 1 / (1 + exp(-x %*% theta))) * x
}
logistic_loss_hessian <- function(data, theta) {
  x <- data[nrow(data), -1]
  prob <- 1 / (1 + exp(-x %*% theta))
  (x %o% x) * c((1 - prob) * prob)
}
result_custom <- fastcpd(
  formula = y ~ . - 1,
  data = data,
  epsilon = 1e-5,
  cost = logistic_loss,
  cost_gradient = logistic_loss_gradient,
  cost_hessian = logistic_loss_hessian
)
result_builtin@cp_set
result_custom@cp_set

```

```

if (requireNamespace("mvtnorm", quietly = TRUE)) {
  set.seed(1)
  n <- 480
  p_true <- 6
  p <- 50
  x <- mvtnorm::rmvnorm(n, rep(0, p), diag(p))
  theta_0 <- rbind(
    runif(p_true, -5, -2),
    runif(p_true, -3, 3),
    runif(p_true, 2, 5),
    runif(p_true, -5, 5)
  )
  theta_0 <- cbind(theta_0, matrix(0, ncol = p - p_true, nrow = 4))
  y <- c(
    x[1:80, ] %*% theta_0[1, ] + rnorm(80, 0, 1),
    x[81:200, ] %*% theta_0[2, ] + rnorm(120, 0, 1),
    x[201:320, ] %*% theta_0[3, ] + rnorm(120, 0, 1),
    x[321:n, ] %*% theta_0[4, ] + rnorm(160, 0, 1)
  )
  small_lasso_data <- cbind.data.frame(y, x)
  result_no_vp <- fastcpd.lasso(
    small_lasso_data,
    beta = "BIC",
    cost_adjustment = NULL,
    pruning_coef = 0
  )
  summary(result_no_vp)
  result_20_vp <- fastcpd.lasso(
    small_lasso_data,
    beta = "BIC",
    cost_adjustment = NULL,
    vanilla_percentage = 0.2,
    pruning_coef = 0
  )
  summary(result_20_vp)
}

```

Description

This S4 class stores the output from [fastcpd\(\)](#) and [fastcpd.family](#). A fastcpd object consist of several slots including the call to [fastcpd\(\)](#), the data used, the family of the model, the change points, the cost values, the residuals, the estimated parameters and a boolean indicating whether the model was fitted with only change points or with change points and parameters, which you can select using @.

Slots

`call` The call of the function.
`data` The data passed to the function.
`order` The order of the time series model.
`family` The family of the model.
`cp_set` The set of change points.
`cost_values` The cost function values for each segment.
`residuals` The residuals of the model with change points. Used only for built-in families.
`thetas` The estimated parameters for each segment. Used only for built-in families.
`cp_only` A boolean indicating whether `fastcpd()` was run to return only the change points or the change points with the estimated parameters and cost values for each segment.

`fastcpd_ar`

Find change points efficiently in AR(p) models

Description

`fastcpd_ar()` and `fastcpd.ar()` are wrapper functions of `fastcpd()` to find change points in AR(p) models. The function is similar to `fastcpd()` except that the data is by default a one-column matrix or univariate vector and thus a formula is not required here.

Usage

```
fastcpd_ar(data, order = 0, ...)
fastcpd.ar(data, order = 0, ...)
```

Arguments

<code>data</code>	A numeric vector, a matrix, a data frame or a time series object.
<code>order</code>	A positive integer specifying the order of the AR model.
<code>...</code>	Other arguments passed to <code>fastcpd()</code> , for example, <code>segment_count</code> . One special argument can be passed here is <code>include.mean</code> , which is a logical value indicating whether the mean should be included in the model. The default value is TRUE.

Value

A `fastcpd` object.

See Also

`fastcpd()`

Examples

```

set.seed(1)
n <- 1000
x <- rep(0, n + 3)
for (i in 1:600) {
  x[i + 3] <- 0.6 * x[i + 2] - 0.2 * x[i + 1] + 0.1 * x[i] + rnorm(1, 0, 3)
}
for (i in 601:1000) {
  x[i + 3] <- 0.3 * x[i + 2] + 0.4 * x[i + 1] + 0.2 * x[i] + rnorm(1, 0, 3)
}
result <- fastcpd.ar(x[3 + seq_len(n)], 3)
summary(result)
plot(result)
set.seed(1)
n <- 1000
x <- rep(0, n + 3)
for (i in 1:600) {
  x[i + 3] <- 0.6 * x[i + 2] - 0.2 * x[i + 1] + 0.1 * x[i] + rnorm(1, 0, 3)
}
for (i in 601:1000) {
  x[i + 3] <- 0.3 * x[i + 2] + 0.4 * x[i + 1] + 0.2 * x[i] + rnorm(1, 0, 3)
}
result <-
  fastcpd.ar(x[3 + seq_len(n)], 3, beta = "MDL", cost_adjustment = "MDL")
summary(result)
plot(result)

```

fastcpd_arima

Find change points efficiently in ARIMA(p, d, q) models

Description

`fastcpd_arima()` and `fastcpd.arima()` are wrapper functions of `fastcpd()` to find change points in ARIMA(p, d, q) models. The function is similar to `fastcpd()` except that the data is by default a one-column matrix or univariate vector and thus a formula is not required here.

Usage

```

fastcpd_arima(data, order = 0, ...)
fastcpd.arima(data, order = 0, ...)

```

Arguments

data	A numeric vector, a matrix, a data frame or a time series object.
order	A vector of length three specifying the order of the ARIMA model.
...	Other arguments passed to <code>fastcpd()</code> , for example, <code>segment_count</code> . One special argument can be passed here is <code>include.mean</code> , which is a logical value indicating whether the mean should be included in the model. The default value is TRUE.

Value

A `fastcpd` object.

See Also

`fastcpd()`

Examples

```
set.seed(1)
n <- 801
w <- rnorm(n + 1, 0, 3)
dx <- rep(0, n + 1)
x <- rep(0, n + 1)
for (i in 1:400) {
  dx[i + 1] <- 0.9 * dx[i] + w[i + 1] - 0.1 * w[i]
  x[i + 1] <- x[i] + dx[i + 1]
}
for (i in 401:n) {
  dx[i + 1] <- -0.6 * dx[i] + w[i + 1] + 0.3 * w[i]
  x[i + 1] <- x[i] + dx[i + 1]
}
result <- fastcpd.arima(
  diff(x[1 + seq_len(n)]),
  c(1, 0, 1),
  segment_count = 3,
  include.mean = FALSE
)
summary(result)
plot(result)
```

`fastcpd_arma`

Find change points efficiently in ARMA(p, q) models

Description

`fastcpd_arma()` and `fastcpd.arma()` are wrapper functions of `fastcpd()` to find change points in ARMA(p, q) models. The function is similar to `fastcpd()` except that the data is by default a one-column matrix or univariate vector and thus a formula is not required here.

Usage

```
fastcpd_arma(data, order = c(0, 0), ...)
fastcpd.arma(data, order = c(0, 0), ...)
```

Arguments

- data A numeric vector, a matrix, a data frame or a time series object.
- order A vector of length two specifying the order of the ARMA model.
- ... Other arguments passed to [fastcpd\(\)](#), for example, segment_count.

Value

A [fastcpd](#) object.

See Also

[fastcpd\(\)](#)

Examples

```
set.seed(1)
n <- 200
w <- rnorm(n + 3, 0, 3)
x <- rep(0, n + 3)
for (i in 1:150) {
  x[i + 3] <- 0.1 * x[i + 2] - 0.3 * x[i + 1] + 0.1 * x[i] +
  0.1 * w[i + 2] + 0.5 * w[i + 1] + w[i + 3]
}
for (i in 151:n) {
  x[i + 3] <- 0.3 * x[i + 2] + 0.1 * x[i + 1] - 0.3 * x[i] -
  0.6 * w[i + 2] - 0.1 * w[i + 1] + w[i + 3]
}
result <- suppressWarnings(
  fastcpd.arma(
    data = x[3 + seq_len(n)],
    order = c(3, 2),
    segment_count = 3,
    lower = c(rep(-1, 3 + 2), 1e-10),
    upper = c(rep(1, 3 + 2), Inf),
    line_search = c(1, 0.1, 1e-2),
    beta = "BIC",
    cost_adjustment = "BIC",
    trim = 0.025
  )
)
summary(result)
plot(result)
```

fastcpd_binomial*Find change points efficiently in logistic regression models***Description**

fastcpd_binomial() and **fastcpd.binomial()** are wrapper functions of **fastcpd()** to find change points in logistic regression models. The function is similar to **fastcpd()** except that the data is by default a matrix or data frame with the response variable as the first column and thus a formula is not required here.

Usage

```
fastcpd_binomial(data, ...)
fastcpd.binomial(data, ...)
```

Arguments

data	A matrix or a data frame with the response variable as the first column.
...	Other arguments passed to fastcpd() , for example, <code>segment_count</code> .

Value

A **fastcpd** object.

See Also

fastcpd()

Examples

```
if (requireNamespace("mvtnorm", quietly = TRUE)) {
  set.seed(1)
  n <- 500
  p <- 4
  x <- mvtnorm::rmvnorm(n, rep(0, p), diag(p))
  theta <- rbind(rnorm(p, 0, 1), rnorm(p, 2, 1))
  y <- c(
    rbinom(300, 1, 1 / (1 + exp(-x[1:300, ] %*% theta[1, ]))),
    rbinom(200, 1, 1 / (1 + exp(-x[301:n, ] %*% theta[2, ])))
  )
  result <- suppressWarnings(fastcpd.binomial(cbind(y, x)))
  summary(result)
  plot(result)
}
```

fastcpd_garch	<i>Find change points efficiently in GARCH(p, q) models</i>
---------------	--

Description

`fastcpd_garch()` and `fastcpd.garch()` are wrapper functions of `fastcpd()` to find change points in GARCH(p, q) models. The function is similar to `fastcpd()` except that the data is by default a one-column matrix or univariate vector and thus a formula is not required here.

Usage

```
fastcpd_garch(data, order = c(0, 0), ...)
fastcpd.garch(data, order = c(0, 0), ...)
```

Arguments

- | | |
|--------------------|--|
| <code>data</code> | A numeric vector, a matrix, a data frame or a time series object. |
| <code>order</code> | A positive integer vector of length two specifying the order of the GARCH model. |
| <code>...</code> | Other arguments passed to <code>fastcpd()</code> , for example, <code>segment_count</code> . |

Value

A `fastcpd` object.

See Also

`fastcpd()`

Examples

```
set.seed(1)
n <- 1501
sigma_2 <- rep(1, n + 1)
x <- rep(0, n + 1)
for (i in seq_len(750)) {
  sigma_2[i + 1] <- 20 + 0.8 * x[i]^2 + 0.1 * sigma_2[i]
  x[i + 1] <- rnorm(1, 0, sqrt(sigma_2[i + 1]))
}
for (i in 751:n) {
  sigma_2[i + 1] <- 1 + 0.1 * x[i]^2 + 0.5 * sigma_2[i]
  x[i + 1] <- rnorm(1, 0, sqrt(sigma_2[i + 1]))
}
result <- suppressWarnings(
  fastcpd.garch(x[-1], c(1, 1), include.mean = FALSE)
)
summary(result)
```

```
plot(result)
```

fastcpd_lasso

Find change points efficiently in penalized linear regression models

Description

fastcpd_lasso() and **fastcpd.lasso()** are wrapper functions of **fastcpd()** to find change points in penalized linear regression models. The function is similar to **fastcpd()** except that the data is by default a matrix or data frame with the response variable as the first column and thus a formula is not required here.

Usage

```
fastcpd_lasso(data, ...)
fastcpd.lasso(data, ...)
```

Arguments

data	A matrix or a data frame with the response variable as the first column.
...	Other arguments passed to fastcpd() , for example, segment_count.

Value

A **fastcpd** object.

See Also

[fastcpd\(\)](#)

Examples

```
if (
  requireNamespace("ggplot2", quietly = TRUE) &&
  requireNamespace("mvtnorm", quietly = TRUE)
) {
  set.seed(1)
  n <- 480
  p_true <- 5
  p <- 50
  x <- mvtnorm::rmvnorm(n, rep(0, p), diag(p))
  theta_0 <- rbind(
    runif(p_true, -5, -2),
    runif(p_true, -3, 3),
    runif(p_true, 2, 5),
    runif(p_true, -5, 5)
  )
```

```

theta_0 <- cbind(theta_0, matrix(0, ncol = p - p_true, nrow = 4))
y <- c(
  x[1:80, ] %*% theta_0[1, ] + rnorm(80, 0, 1),
  x[81:200, ] %*% theta_0[2, ] + rnorm(120, 0, 1),
  x[201:320, ] %*% theta_0[3, ] + rnorm(120, 0, 1),
  x[321:n, ] %*% theta_0[4, ] + rnorm(160, 0, 1)
)
result <- fastcpd.lasso(
  cbind(y, x),
  multiple_epochs = function(segment_length) if (segment_length < 30) 1 else 0
)
summary(result)
plot(result)

# Combine estimated thetas with true parameters
thetas <- result@thetas
thetas <- cbind.data.frame(thetas, t(theta_0))
names(thetas) <- c(
  "segment 1", "segment 2", "segment 3", "segment 4",
  "segment 1 truth", "segment 2 truth", "segment 3 truth", "segment 4 truth"
)
thetas$coordinate <- c(seq_len(p_true), rep("rest", p - p_true))

# Melt the data frame using base R (i.e., convert from wide to long format)
data_cols <- setdiff(names(thetas), "coordinate")
molten <- data.frame(
  coordinate = rep(thetas$coordinate, times = length(data_cols)),
  variable = rep(data_cols, each = nrow(thetas)),
  value = as.vector(as.matrix(thetas[, data_cols]))
)

# Remove the "segment " and " truth" parts to extract the segment number
molten$segment <- gsub("segment ", "", molten$variable)
molten$segment <- gsub(" truth", "", molten$segment)

# Compute height: the numeric value of the segment plus an offset for truth values
molten$height <- as.numeric(gsub("segment.*", "", molten$segment)) +
  0.2 * as.numeric(grepl("truth", molten$variable))

# Create a parameter indicator based on whether the variable corresponds to truth or estimation
molten$parameter <- ifelse(grepl("truth", molten$variable), "truth", "estimated")

p <- ggplot2::ggplot() +
  ggplot2::geom_point(
    data = molten,
    ggplot2::aes(
      x = value, y = height, shape = coordinate, color = parameter
    ),
    size = 4
  ) +
  ggplot2::ylim(0.8, 4.4) +
  ggplot2::ylab("segment") +
  ggplot2::theme_bw()

```

```
print(p)
}
```

fastcpd_lm*Find change points efficiently in linear regression models***Description**

fastcpd_lm() and **fastcpd.lm()** are wrapper functions of **fastcpd()** to find change points in linear regression models. The function is similar to **fastcpd()** except that the data is by default a matrix or data frame with the response variable as the first column and thus a formula is not required here.

Usage

```
fastcpd_lm(data, ...)
fastcpd.lm(data, ...)
```

Arguments

data	A matrix or a data frame with the response variable as the first column.
...	Other arguments passed to fastcpd() , for example, <code>segment_count</code> .

Value

A **fastcpd** object.

See Also

[fastcpd\(\)](#)

Examples

```
if (requireNamespace("mvtnorm", quietly = TRUE)) {
  set.seed(1)
  n <- 300
  p <- 4
  x <- mvtnorm::rmvnorm(n, rep(0, p), diag(p))
  theta_0 <- rbind(c(1, 3.2, -1, 0), c(-1, -0.5, 2.5, -2), c(0.8, 0, 1, 2))
  y <- c(
    x[1:100, ] %*% theta_0[1, ] + rnorm(100, 0, 3),
    x[101:200, ] %*% theta_0[2, ] + rnorm(100, 0, 3),
    x[201:n, ] %*% theta_0[3, ] + rnorm(100, 0, 3)
  )
  result_lm <- fastcpd.lm(cbind(y, x))
  summary(result_lm)
  plot(result_lm)
```

```

}

if (requireNamespace("mvtnorm", quietly = TRUE)) {
  set.seed(1)
  n <- 600
  p <- 4
  d <- 2
  x <- mvtnorm::rmvnorm(n, rep(0, p), diag(p))
  theta_1 <- matrix(runif(8, -3, -1), nrow = p)
  theta_2 <- matrix(runif(8, -1, 3), nrow = p)
  y <- rbind(
    x[1:350, ] %*% theta_1 + mvtnorm::rmvnorm(350, rep(0, d), 3 * diag(d)),
    x[351:n, ] %*% theta_2 + mvtnorm::rmvnorm(250, rep(0, d), 3 * diag(d))
  )
  result_mlm <- fastcpd.lm(cbind.data.frame(y = y, x = x), p.response = 2)
  summary(result_mlm)
}

```

fastcpd_mean*Find change points efficiently in mean change models***Description**

`fastcpd_mean()` and `fastcpd.mean()` are wrapper functions of `fastcpd()` to find the mean change. The function is similar to `fastcpd()` except that the data is by default a matrix or data frame or a vector with each row / element as an observation and thus a formula is not required here.

Usage

```
fastcpd_mean(data, ...)
fastcpd.mean(data, ...)
```

Arguments

<code>data</code>	A matrix, a data frame or a vector.
<code>...</code>	Other arguments passed to <code>fastcpd()</code> , for example, <code>segment_count</code> .

Value

A `fastcpd` object.

See Also

`fastcpd()`

Examples

```

set.seed(1)
data <- matrix(c(
  rnorm(300, mean = 0, sd = 10),
  rnorm(400, mean = 50, sd = 10),
  rnorm(300, mean = 2, sd = 10)
))
system.time(result <- fastcpd.mean(data))
summary(result)
plot(result)
set.seed(1)
p <- 3
data <- rbind(
  matrix(rnorm(p * 3e+5, mean = 0, sd = 10), ncol = p),
  matrix(rnorm(p * 4e+5, mean = 50, sd = 10), ncol = p),
  matrix(rnorm(p * 3e+5, mean = 2, sd = 10), ncol = p)
)
system.time(result <- fastcpd.mean(data, r.progress = FALSE, cp_only = TRUE))
summary(result)

```

fastcpd_meanvariance *Find change points efficiently in mean variance change models*

Description

fastcpd_meanvariance(), **fastcpd.meanvariance()**, **fastcpd_mv()**, **fastcpd.mv()** are wrapper functions of **fastcpd()** to find the meanvariance change. The function is similar to **fastcpd()** except that the data is by default a matrix or data frame or a vector with each row / element as an observation and thus a formula is not required here.

Usage

```

fastcpd_meanvariance(data, ...)
fastcpd.meanvariance(data, ...)
fastcpd_mv(data, ...)
fastcpd.mv(data, ...)

```

Arguments

data	A matrix, a data frame or a vector.
...	Other arguments passed to fastcpd() , for example, <code>segment_count</code> .

Value

A **fastcpd** object.

See Also

[fastcpd\(\)](#)

Examples

```
set.seed(1)
data <- c(
  rnorm(3000, 0, 1),
  rnorm(1000, 10, 1),
  rnorm(3000, 10, 20),
  rnorm(1000, 0, 1)
)
system.time(result <- fastcpd.mv(data))
summary(result)
plot(result)
set.seed(1)
p <- 3
data <- if (requireNamespace("mvtnorm", quietly = TRUE)) {
  rbind(
    mvtnorm::rmvnorm(2e+5, mean = rep(0, p), sigma = diag(1, p)),
    mvtnorm::rmvnorm(1e+5, mean = rep(50, p), sigma = diag(1, p)),
    mvtnorm::rmvnorm(2e+5, mean = rep(0, p), sigma = diag(100, p)),
    mvtnorm::rmvnorm(2e+5, mean = rep(0, p), sigma = diag(1, p)),
    mvtnorm::rmvnorm(1e+5, mean = rep(50, p), sigma = diag(1, p)),
    mvtnorm::rmvnorm(2e+5, mean = rep(50, p), sigma = diag(100, p))
  )
} else {
  rbind(
    matrix(rnorm(p * 2e+5, mean = 0, sd = 1), ncol = p),
    matrix(rnorm(p * 1e+5, mean = 50, sd = 1), ncol = p),
    matrix(rnorm(p * 2e+5, mean = 0, sd = 10), ncol = p),
    matrix(rnorm(p * 2e+5, mean = 0, sd = 1), ncol = p),
    matrix(rnorm(p * 1e+5, mean = 50, sd = 1), ncol = p),
    matrix(rnorm(p * 2e+5, mean = 50, sd = 10), ncol = p)
  )
}
system.time(result <- fastcpd.mv(data))
summary(result)
result@thetas[seq_len(p), ]
lapply(result@thetas[seq_len(p^2) + p, ], function(thetas) matrix(thetas, p))
```

fastcpd_poisson

*Find change points efficiently in Poisson regression models***Description**

[fastcpd_poisson\(\)](#) and [fastcpd.poisson\(\)](#) are wrapper functions of [fastcpd\(\)](#) to find change points in Poisson regression models. The function is similar to [fastcpd\(\)](#) except that the data is by default a matrix or data frame with the response variable as the first column and thus a formula is not required here.

Usage

```
fastcpd_poisson(data, ...)
fastcpd.poisson(data, ...)
```

Arguments

- `data` A matrix or a data frame with the response variable as the first column.
`...` Other arguments passed to [fastcpd\(\)](#), for example, `segment_count`.

Value

A [fastcpd](#) object.

See Also

[fastcpd\(\)](#)

Examples

```
if (requireNamespace("mvtnorm", quietly = TRUE)) {
  set.seed(1)
  n <- 1100
  p <- 3
  x <- mvtnorm::rmvnorm(n, rep(0, p), diag(p))
  delta <- rnorm(p)
  theta_0 <- c(1, 0.3, -1)
  y <- c(
    rpois(500, exp(x[1:500, ] %*% theta_0)),
    rpois(300, exp(x[501:800, ] %*% (theta_0 + delta))),
    rpois(200, exp(x[801:1000, ] %*% theta_0)),
    rpois(100, exp(x[1001:1100, ] %*% (theta_0 - delta)))
  )
  result <- fastcpd.poisson(cbind(y, x))
  summary(result)
  plot(result)
}
```

fastcpd_ts

Find change points efficiently in time series data

Description

[fastcpd_ts\(\)](#) and [fastcpd.ts\(\)](#) are wrapper functions for [fastcpd\(\)](#) to find change points in time series data. The function is similar to [fastcpd\(\)](#) except that the data is a time series and the family is one of "ar", "var", "arma", "arima" or "garch".

Usage

```
fastcpd_ts(data, family = NULL, order = c(0, 0, 0), ...)
fastcpd.ts(data, family = NULL, order = c(0, 0, 0), ...)
```

Arguments

data	A numeric vector, a matrix, a data frame or a time series object.
family	A character string specifying the family of the time series. The value should be one of "ar", "var", "arima" or "garch".
order	A positive integer or a vector of length less than four specifying the order of the time series. Possible combinations with family are:
	<ul style="list-style-type: none"> • "ar", NUMERIC(1): AR(p) model using linear regression. • "var", NUMERIC(1): VAR(p) model using linear regression. • "arima", NUMERIC(3): ARIMA(p, d, q) model using stats:::arima(). • "garch", NUMERIC(2): GARCH(p, q) model.
...	Other arguments passed to fastcpd() , for example, segment_count. One special argument can be passed here is include.mean, which is a logical value indicating whether the mean should be included in the model. The default value is TRUE.

Value

A [fastcpd](#) object.

See Also

[fastcpd\(\)](#)

Examples

```
# Set seed for reproducibility
set.seed(1)

# 1. Define Parameters
n <- 500          # Total length of the time series
cp1 <- 200        # First change point at time 200
cp2 <- 350        # Second change point at time 350

# Define MA(2) coefficients for each segment ensuring invertibility
# MA coefficients affect invertibility; to ensure invertibility, the roots of
# the MA polynomial should lie outside the unit circle.

# Segment 1: Time 1 to cp1
theta1_1 <- 0.5
theta2_1 <- -0.3

# Segment 2: Time (cp1+1) to cp2
theta1_2 <- -0.4
```

```

theta2_2 <- 0.25

# Segment 3: Time (cp2+1) to n
theta1_3 <- 0.6
theta2_3 <- -0.35

# Function to check invertibility for MA(2)
is_invertible_ma2 <- function(theta1, theta2) {
  # The MA(2) polynomial is: 1 + theta1*z + theta2*z^2 = 0
  # Compute the roots of the polynomial
  roots <- polyroot(c(1, theta1, theta2))
  # Invertible if all roots lie outside the unit circle
  all(Mod(roots) > 1)
}

# Verify invertibility for each segment
stopifnot(is_invertible_ma2(theta1_1, theta2_1))
stopifnot(is_invertible_ma2(theta1_2, theta2_2))
stopifnot(is_invertible_ma2(theta1_3, theta2_3))

# 2. Simulate White Noise
e <- rnorm(n + 2, mean = 0, sd = 1) # Extra terms to handle lag

# 3. Initialize the Time Series
y <- numeric(n + 2) # Extra terms for initial lags (y[1], y[2] are zero)

# 4. Apply MA(2) Model with Change Points
for (t in 3:(n + 2)) { # Start from 3 to have enough lags for MA(2)
  # Determine current segment
  current_time <- t - 2 # Adjust for the extra lags
  if (current_time <= cp1) {
    theta <- c(theta1_1, theta2_1)
  } else if (current_time <= cp2) {
    theta <- c(theta1_2, theta2_2)
  } else {
    theta <- c(theta1_3, theta2_3)
  }

  # Compute MA(2) value
  y[t] <- e[t] + theta[1] * e[t - 1] + theta[2] * e[t - 2]
}

# Remove the initial extra terms
y <- y[3:(n + 2)]
time <- 1:n

# Function to get roots data for plotting
get_roots_data <- function(theta1, theta2, segment) {
  roots <- polyroot(c(1, theta1, theta2))
  data.frame(
    Re = Re(roots),
    Im = Im(roots),
    Distance = Mod(roots),
  )
}

```

```

    Segment = segment
)
}

roots_segment1 <- get_roots_data(theta1_1, theta2_1, "Segment 1")
roots_segment2 <- get_roots_data(theta1_2, theta2_2, "Segment 2")
roots_segment3 <- get_roots_data(theta1_3, theta2_3, "Segment 3")

(roots_data <- rbind(roots_segment1, roots_segment2, roots_segment3))

result <- fastcpd.ts(
  y,
  "arma",
  c(0, 2),
  lower = c(-2, -2, 1e-10),
  upper = c(2, 2, Inf),
  line_search = c(1, 0.1, 1e-2),
  trim = 0.04
)
summary(result)
plot(result)

```

fastcpd_var*Find change points efficiently in VAR(p) models***Description**

`fastcpd_var()` and `fastcpd.var()` are wrapper functions of `fastcpd_ts()` to find change points in VAR(p) models. The function is similar to `fastcpd_ts()` except that the data is by default a matrix with row as an observation and thus a formula is not required here.

Usage

```
fastcpd_var(data, order = 0, ...)
fastcpd.var(data, order = 0, ...)
```

Arguments

- | | |
|--------------------|--|
| <code>data</code> | A matrix, a data frame or a time series object. |
| <code>order</code> | A positive integer specifying the order of the VAR model. |
| <code>...</code> | Other arguments passed to <code>fastcpd()</code> , for example, <code>segment_count</code> . |

Value

A `fastcpd` object.

See Also[fastcpd\(\)](#)**Examples**

```
set.seed(1)
n <- 300
p <- 2
theta_1 <- matrix(c(-0.3, 0.6, -0.5, 0.4, 0.2, 0.2, 0.2, -0.2), nrow = p)
theta_2 <- matrix(c(0.3, -0.4, 0.1, -0.5, -0.5, -0.2, -0.5, 0.2), nrow = p)
x <- matrix(0, n + 2, p)
for (i in 1:200) {
  x[i + 2, ] <- theta_1 %*% c(x[i + 1, ], x[i, ]) + rnorm(p, 0, 1)
}
for (i in 201:n) {
  x[i + 2, ] <- theta_2 %*% c(x[i + 1, ], x[i, ]) + rnorm(p, 0, 1)
}
result <- fastcpd.var(x, 2)
summary(result)
```

fastcpd_variance*Find change points efficiently in variance change models***Description**

fastcpd_variance() and **fastcpd.variance()** are wrapper functions of **fastcpd()** to find the variance change. The function is similar to **fastcpd()** except that the data is by default a matrix or data frame or a vector with each row / element as an observation and thus a formula is not required here.

Usage**fastcpd_variance(data, ...)****fastcpd.variance(data, ...)****Arguments**

data A matrix, a data frame or a vector.

... Other arguments passed to **fastcpd()**, for example, `segment_count`.**Value**A **fastcpd** object.**See Also**[fastcpd\(\)](#)

Examples

```

set.seed(1)
data <- c(rnorm(300, 0, 1), rnorm(400, 0, 10), rnorm(300, 0, 1))
system.time(result <- fastcpd.variance(data))
summary(result)
plot(result)
if (requireNamespace("mvtnorm", quietly = TRUE)) {
  set.seed(1)
  p <- 3
  data <- rbind(
    mvtnorm::rmvnorm(
      3e+5, rep(0, p), crossprod(matrix(runif(p^2) * 2 - 1, p)))
    ),
    mvtnorm::rmvnorm(
      4e+5, rep(0, p), crossprod(matrix(runif(p^2) * 2 - 1, p)))
    ),
    mvtnorm::rmvnorm(
      3e+5, rep(0, p), crossprod(matrix(runif(p^2) * 2 - 1, p)))
    )
  )
  result_time <- system.time(
    result <- fastcpd.variance(data, r.progress = FALSE, cp_only = TRUE)
  )
  print(result_time)
  summary(result)
}

```

occupancy

Occupancy Detection Data Set

Description

Data set for binary classification of room occupancy from temperature, humidity, light and CO₂ measurements. Ground-truth occupancy was obtained from time stamped pictures that were taken every minute.

Usage

```
occupancy
```

Format

A data frame with 9752 rows and 7 variables:

date Character in the format "YYYY-MM-DD hh:mm:ss" from 2015-02-11 14:48:00 to 2015-02-18 09:19:00

Temperature Temperature in Celsius

Humidity Humidity

Light Light

CO2 CO2

HumidityRatio Humidity Ratio

Occupancy Binary variable with values 0 (unoccupied) and 1

Source

<<https://github.com/LuisM78/Occupancy-detection-data>>

plot.fastcpd

Plot the data and the change points for a [fastcpd](#) object

Description

Plot the data and the change points for a [fastcpd](#) object

Usage

```
## S3 method for class 'fastcpd'
plot(
  x,
  color_max_count = Inf,
  data_point_alpha = 0.8,
  data_point_linewidth = 0.5,
  data_point_size = 1,
  legend_position = "none",
  panel_background = ggplot2::element_blank(),
  panel_border = ggplot2::element_rect(fill = NA, colour = "grey20"),
  panel_grid_major = ggplot2::element_line(colour = "grey98"),
  panel_grid_minor = ggplot2::element_line(colour = "grey98"),
  segment_separator_alpha = 0.8,
  segment_separator_color = "grey",
  segment_separator_linetype = "dashed",
  strip_background = ggplot2::element_rect(fill = "grey85", colour = "grey20"),
  xlab = NULL,
  ylab = NULL,
  ...
)

## S4 method for signature 'fastcpd,missing'
plot(
  x,
  color_max_count = Inf,
  data_point_alpha = 0.8,
  data_point_linewidth = 0.5,
  data_point_size = 1,
```

```
legend_position = "none",
panel_background = ggplot2::element_blank(),
panel_border = ggplot2::element_rect(fill = NA, colour = "grey20"),
panel_grid_major = ggplot2::element_line(colour = "grey98"),
panel_grid_minor = ggplot2::element_line(colour = "grey98"),
segment_separator_alpha = 0.8,
segment_separator_color = "grey",
segment_separator_linetype = "dashed",
strip_background = ggplot2::element_rect(fill = "grey85", colour = "grey20"),
xlab = NULL,
ylab = NULL,
...
)
```

Arguments

x	A fastcpd object.
color_max_count	Maximum number of colors to use for the plotting of segments.
data_point_alpha	Alpha of the data points.
data_point_linewidth	Linewidth of the data points.
data_point_size	Size of the data points.
legend_position	Position of the legend.
panel_background	Background of the panel.
panel_border	Border of the panel.
panel_grid_major	Major grid lines of the panel.
panel_grid_minor	Minor grid lines of the panel.
segment_separator_alpha	Alpha of the segment separator lines.
segment_separator_color	Color of the segment separator lines.
segment_separator_linetype	Linetype of the segment separator lines.
strip_background	Background of the strip.
xlab	Label for the x-axis.
ylab	Label for the y-axis.
...	Ignored.

Value

No return value, called for plotting.

Examples

```
if (requireNamespace("mvtnorm", quietly = TRUE)) {
  set.seed(1)
  p <- 1
  x <- mvtnorm::rmvnorm(300, rep(0, p), diag(p))
  theta_0 <- matrix(c(1, -1, 0.5))
  y <- c(
    x[1:100, ] * theta_0[1, ] + rnorm(100, 0, 1),
    x[101:200, ] * theta_0[2, ] + rnorm(100, 0, 1),
    x[201:300, ] * theta_0[3, ] + rnorm(100, 0, 1)
  )
  result <- fastcpd.lm(cbind(y, x))
  summary(result)
  plot(result)
}
```

print.fastcpd

Print the call and the change points for a [fastcpd](#) object

Description

Print the call and the change points for a [fastcpd](#) object

Usage

```
## S3 method for class 'fastcpd'
print(x, ...)

## S4 method for signature 'fastcpd'
print(x, ...)
```

Arguments

x	A fastcpd object.
...	Ignored.

Value

Return a (temporarily) invisible copy of the [fastcpd](#) object. Called primarily for printing the change points in the model.

show.fastcpd	<i>Show the available methods for a fastcpd object</i>
--------------	--

Description

Show the available methods for a [fastcpd](#) object

Usage

```
## S3 method for class 'fastcpd'  
show(object)  
  
## S4 method for signature 'fastcpd'  
show(object)
```

Arguments

object A [fastcpd](#) object.

Value

No return value, called for showing a list of available methods for a [fastcpd](#) object.

summary.fastcpd	<i>Show the summary of a fastcpd object</i>
-----------------	---

Description

Show the summary of a [fastcpd](#) object

Usage

```
## S3 method for class 'fastcpd'  
summary(object, ...)  
  
## S4 method for signature 'fastcpd'  
summary(object, ...)
```

Arguments

object A [fastcpd](#) object.
... Ignored.

Value

Return a (temporarily) invisible copy of the [fastcpd](#) object. Called primarily for printing the summary of the model including the call, the change points, the cost values and the estimated parameters.

transcriptome

Transcription Profiling of 57 Human Bladder Carcinoma Samples

Description

Transcriptome analysis of 57 bladder carcinomas on Affymetrix HG-U95A and HG-U95Av2 microarrays

Usage

transcriptome

Format

A data frame with 2215 rows and 43 variables:

- 3** Individual 3
- 4** Individual 4
- 5** Individual 5
- 6** Individual 6
- 7** Individual 7
- 8** Individual 8
- 9** Individual 9
- 10** Individual 10
- 14** Individual 14
- 15** Individual 15
- 16** Individual 16
- 17** Individual 17
- 18** Individual 18
- 19** Individual 19
- 21** Individual 21
- 22** Individual 22
- 24** Individual 24
- 26** Individual 26
- 28** Individual 28
- 30** Individual 30
- 31** Individual 31
- 33** Individual 33
- 34** Individual 34
- 35** Individual 35

36 Individual 36
37 Individual 37
38 Individual 38
39 Individual 39
40 Individual 40
41 Individual 41
42 Individual 42
43 Individual 43
44 Individual 44
45 Individual 45
46 Individual 46
47 Individual 47
48 Individual 48
49 Individual 49
50 Individual 50
51 Individual 51
53 Individual 53
54 Individual 54
57 Individual 57

Source

<<https://www.ebi.ac.uk/biostudies/arrayexpress/studies/E-TABM-147>>
<<https://github.com/cran/ecp/tree/master/data>>

Examples

```
if (requireNamespace("ggplot2", quietly = TRUE)) {  
  result <- fastcpd.mean(transcriptome$"10", trim = 0.005)  
  summary(result)  
  plot(result)  
  
  result_all <- fastcpd.mean(  
    transcriptome,  
    beta = (ncol(transcriptome) + 1) * log(nrow(transcriptome)) / 2 * 5,  
    trim = 0  
  )  
  
  plots <- lapply(  
    seq_len(ncol(transcriptome)), function(i) {  
      ggplot2::ggplot(  
        data = data.frame(  
          x = seq_along(transcriptome[, i]), y = transcriptome[, i]  
        ),  
        ggplot2::aes(x = x, y = y)
```

```

) +
  ggplot2::geom_line(color = "steelblue") +
  ggplot2::geom_vline(
    xintercept = result_all@cp_set,
    color = "red",
    linetype = "dotted",
    linewidth = 0.5,
    alpha = 0.7
  ) +
  ggplot2::theme_void()
}
)

if (requireNamespace("gridExtra", quietly = TRUE)) {
  gridExtra::grid.arrange(grobs = plots, ncol = 1, nrow = ncol(transcriptome))
}
}

```

uk_seatbelts*UK Seatbelts Data***Description**

Road Casualties in Great Britain 1969–84.

Usage

`uk_seatbelts`

Format

`uk_seatbelts` is a multiple time series, with columns

DriversKilled car drivers killed.

front front-seat passengers killed or seriously injured.

rear rear-seat passengers killed or seriously injured.

kms distance driven.

PetrolPrice petrol price.

VanKilled number of van ('light goods vehicle') drivers.

law 0/1: was the law in effect that month?

Source

R package **datasets**

Examples

```

if (requireNamespace("ggplot2", quietly = TRUE)) {
  result_ar <- fastcpd.ar(diff(uk_seatbelts[, "drivers"], 12), 1, beta = "BIC")
  summary(result_ar)
  plot(result_ar)

  result_lm <- suppressMessages(fastcpd.lm(
    diff(uk_seatbelts[, c("drivers", "kms", "PetrolPrice", "law")]), lag = 12
  ))

  # Compute change point dates:
  cp_dates <- as.POSIXlt(as.Date("1969-01-01", format = "%Y-%m-%d"))
  cp_dates$mon <- cp_dates$mon + (1 + result_lm@cp_set + 12)
  cp_dates <- as.Date(cp_dates)

  # Convert the time series to Date objects:
  # For a monthly ts object, extract year and month manually.
  time_vals <- time(uk_seatbelts)
  years <- floor(time_vals)
  months <- round((time_vals - years) * 12 + 1)
  dates <- as.Date(paste(years, months, "01", sep = "-"), format = "%Y-%m-%d")

  # Prepare the data frame for plotting
  # 'color' is defined similarly to the original code.
  uk_seatbelts_df <- data.frame(
    dates = dates,
    drivers = as.numeric(uk_seatbelts[, "drivers"]),
    color = as.factor((dates < cp_dates[1]) + (dates < cp_dates[2]))
  )

  p <- ggplot2::ggplot(
    data = uk_seatbelts_df,
    ggplot2::aes(x = dates, y = drivers, color = color)
  ) +
    ggplot2::geom_line() +
    ggplot2::geom_vline(
      xintercept = cp_dates,
      linetype = "dashed",
      color = "red"
    ) +
    ggplot2::scale_x_date(date_labels = "%b %Y", date_breaks = "1 year") +
    ggplot2::annotate(
      "text",
      x = cp_dates,
      y = 1025,
      label = format(cp_dates, "%b %Y"),
      color = "blue"
    ) +
    ggplot2::theme_bw() +
    ggplot2::theme(legend.position = "none")
  print(p)
}

```

variance_arma*Variance estimation for ARMA model with change points*

Description

Estimate the variance for each block and then take the average.

Usage

```
variance_arma(data, p, q, max_order = p * q)

variance.arma(data, p, q, max_order = p * q)
```

Arguments

data	A one-column matrix or a vector.
p	The order of the autoregressive part.
q	The order of the moving average part.
max_order	The maximum order of the AR model to consider.

Value

A numeric value representing the variance.

Examples

```
set.seed(1)
n <- 300
w <- rnorm(n + 3, 0, 10)
x <- rep(0, n + 3)
for (i in 1:200) {
  x[i + 3] <- 0.1 * x[i + 2] - 0.3 * x[i + 1] + 0.1 * x[i] +
  0.1 * w[i + 2] + 0.5 * w[i + 1] + w[i + 3]
}
for (i in 201:n) {
  x[i + 3] <- 0.3 * x[i + 2] + 0.1 * x[i + 1] - 0.3 * x[i] -
  0.6 * w[i + 2] - 0.1 * w[i + 1] + w[i + 3]
}
(result <- variance.arma(x[-seq_len(3)], p = 3, q = 2))
```

variance_lm

Variance estimation for linear models with change points

Description

Estimate the variance for each block and then take the average.

Usage

```
variance_lm(data, d = 1, block_size = ncol(data) - d + 1, outlier_iqr = Inf)

variance.lm(data, d = 1, block_size = ncol(data) - d + 1, outlier_iqr = Inf)
```

Arguments

data	A matrix or a data frame with the response variable as the first column.
d	The dimension of the response variable.
block_size	The size of the blocks to use for variance estimation.
outlier_iqr	The number of interquartile ranges to use as a threshold for outlier detection.

Value

A numeric value representing the variance.

Examples

```
if (requireNamespace("mvtnorm", quietly = TRUE)) {
  set.seed(1)
  n <- 300
  p <- 4
  x <- mvtnorm::rmvnorm(n, rep(0, p), diag(p))
  theta <- rbind(c(1, 3.2, -1, 0), c(-1, -0.5, 2.5, -2), c(0.8, 0, 1, 2))
  y <- c(
    x[1:100, ] %*% theta[1, ] + rnorm(100, 0, 3),
    x[101:200, ] %*% theta[2, ] + rnorm(100, 0, 3),
    x[201:n, ] %*% theta[3, ] + rnorm(100, 0, 3)
  )
  (sigma2 <- variance.lm(cbind(y, x)))

  set.seed(1)
  n <- 300
  p <- 4
  d <- 2
  x <- mvtnorm::rmvnorm(n, rep(0, p), diag(p))
  theta <- cbind(c(1, 3.2, -1, 0), c(-1, -0.5, 2.5, -2), c(0.8, 0, 1, 2))
  theta <- cbind(theta, theta)
  y <- rbind(
    x[1:100, ] %*% theta[, 1:2] +
```

```

  mvtnorm::rmvnorm(100, rep(0, d), diag(3, d)),
  x[101:200, ] %*% theta[, 3:4] +
  mvtnorm::rmvnorm(100, rep(0, d), diag(3, d)),
  x[201:n, ] %*% theta[, 5:6] +
  mvtnorm::rmvnorm(100, rep(0, d), diag(3, d))
)
(sigma <- variance.lm(cbind(y, x), d = d))
}

```

variance_mean*Variance estimation for mean change models***Description**

Implement Rice estimator for variance in mean change models.

Usage

```

variance_mean(data)

variance.mean(data)

```

Arguments

data A matrix or a data frame with data points as each row.

Value

A matrix representing the variance-covariance matrix or a numeric value representing the variance.

Examples

```

if (requireNamespace("mvtnorm", quietly = TRUE)) {
  set.seed(1)
  p <- 3
  data <- rbind(
    mvtnorm::rmvnorm(300, mean = rep(0, p), sigma = diag(100, p)),
    mvtnorm::rmvnorm(400, mean = rep(50, p), sigma = diag(100, p)),
    mvtnorm::rmvnorm(300, mean = rep(2, p), sigma = diag(100, p))
  )
  (sigma <- variance.mean(data))
}

```

variance_median	<i>Variance estimation for median change models</i>
-----------------	---

Description

Implement Rice estimator.

Usage

```
variance_median(data)  
variance.median(data)
```

Arguments

data A vector of data points.

Value

A numeric value representing the variance.

Examples

```
(sigma2 <- variance.median(well_log))
```

well_log	<i>Well-log Dataset from Numerical Bayesian Methods Applied to Signal Processing</i>
----------	--

Description

This is the well-known well-log dataset used in many changepoint papers obtained from Alan Turing Institute GitHub repository and licensed under the MIT license.

Usage

```
well_log
```

Format

A Time-Series of length 4050.

Source

<<https://github.com/alan-turing-institute/TCPD>>

Examples

```
result <- fastcpd.mean(well_log, trim = 0.001)
summary(result)
plot(result)

if (requireNamespace("matrixStats", quietly = TRUE)) {
  sigma2 <- variance.median(well_log)
  median_loss <- function(data) {
    sum(abs(data - matrixStats::colMedians(data))) / sqrt(sigma2) / 2
  }
  result <- fastcpd(
    formula = ~ x - 1,
    data = cbind.data.frame(x = well_log),
    cost = median_loss,
    trim = 0.002
  )
  summary(result)

  segment_starts <- c(1, result@cp_set)
  segment_ends <- c(result@cp_set - 1, length(well_log))
  residual <- NULL
  for (segment_index in seq_along(segment_starts)) {
    segment <-
      well_log[segment_starts[segment_index]:segment_ends[segment_index]]
    residual <- c(residual, segment - median(segment))
  }

  result@residuals <- matrix(residual)
  result@data <- data.frame(x = c(well_log))
  plot(result)
}
```

Index

* datasets
 bitcoin, 3
 occupancy, 29
 transcriptome, 34
 uk_seatbelts, 36
 well_log, 41

 bitcoin, 3

 fastcpd, 4, 4, 8, 12, 14–18, 20–22, 24, 25, 27, 28, 30–33
 fastcpd(), 4, 6, 11–18, 20–25, 27, 28
 fastcpd-class, 11
 fastcpd.ar (fastcpd_ar), 12
 fastcpd.ar(), 12
 fastcpd.arima (fastcpd_arima), 13
 fastcpd.arima(), 13
 fastcpd.arma (fastcpd_arma), 14
 fastcpd.arma(), 14
 fastcpd.binomial (fastcpd_binomial), 16
 fastcpd.binomial(), 16
 fastcpd.family, 5, 8, 11
 fastcpd.garch (fastcpd_garch), 17
 fastcpd.garch(), 17
 fastcpd.lasso (fastcpd_lasso), 18
 fastcpd.lasso(), 18
 fastcpd.lm (fastcpd_lm), 20
 fastcpd.lm(), 20
 fastcpd.mean (fastcpd_mean), 21
 fastcpd.mean(), 21
 fastcpd.meanvariance
 (fastcpd_meanvariance), 22
 fastcpd.meanvariance(), 22
 fastcpd.mv (fastcpd_meanvariance), 22
 fastcpd.mv(), 22
 fastcpd.poisson (fastcpd_poisson), 23
 fastcpd.poisson(), 23
 fastcpd.ts (fastcpd_ts), 24
 fastcpd.ts(), 24
 fastcpd.var (fastcpd_var), 27
 fastcpd.var(), 27
 fastcpd.variance (fastcpd_variance), 28
 fastcpd.variance(), 28
 fastcpd_ar, 12
 fastcpd_ar(), 12
 fastcpd_arima, 13
 fastcpd_arima(), 13
 fastcpd_arma, 14
 fastcpd_arma(), 14
 fastcpd_binomial, 16
 fastcpd_binomial(), 16
 fastcpd_garch, 17
 fastcpd_garch(), 17
 fastcpd_lasso, 18
 fastcpd_lasso(), 18
 fastcpd_lm, 20
 fastcpd_lm(), 20
 fastcpd_mean, 21
 fastcpd_mean(), 21
 fastcpd_meanvariance, 22
 fastcpd_meanvariance(), 22
 fastcpd_mv (fastcpd_meanvariance), 22
 fastcpd_mv(), 22
 fastcpd_poisson, 23
 fastcpd_poisson(), 23
 fastcpd_ts, 24
 fastcpd_ts(), 24, 27
 fastcpd_var, 27
 fastcpd_var(), 27
 fastcpd_variance, 28
 fastcpd_variance(), 28
 lm(), 5
 occupancy, 29
 plot, fastcpd, missing-method
 (plot.fastcpd), 30
 plot.fastcpd, 30
 plot.fastcpd(), 8

print, fastcpd-method (print.fastcpd), 32
print.fastcpd, 32

show, fastcpd-method (show.fastcpd), 33
show.fastcpd, 33

stats::arima(), 25

summary, fastcpd-method
(summary.fastcpd), 33

summary.fastcpd, 33

summary.fastcpd(), 8

transcriptome, 34

uk_seatbelts, 36

variance.arma (variance_arma), 38
variance.lm (variance_lm), 39

variance.mean (variance_mean), 40

variance.median (variance_median), 41

variance_arma, 38

variance_lm, 39

variance_mean, 40

variance_median, 41

well_log, 41