

Package ‘hpa’

August 24, 2020

Type Package

Title Distributions Hermite Polynomial Approximation

Version 1.1.2

Date 2020-08-24

Author Potanin Bogdan

Maintainer Potanin Bogdan <bogdanpotanin@gmail.com>

Description Multivariate conditional and marginal densities, moments, cumulative distribution functions as well as binary choice and sample selection models based on hermite polynomial approximation which was proposed and described by A. Gallant and D. W. Ny-chka (1987) <doi:10.2307/1913241>.

License GPL-3

Imports Rcpp (>= 1.0.4), RcppParallel (>= 5.0.0)

LinkingTo Rcpp, RcppArmadillo, RcppParallel

RoxygenNote 7.1.0

Encoding UTF-8

Suggests ggplot2, mvtnorm, titanic, sampleSelection, GA (>= 3.2)

NeedsCompilation yes

SystemRequirements GNU make

Repository CRAN

Date/Publication 2020-08-24 11:00:02 UTC

R topics documented:

AIC.hpaBinary	3
AIC.hpaML	3
AIC.hpaSelection	4
AIC_hpaBinary	4
AIC_hpaML	5
AIC_hpaSelection	5
dhp	6

dhpDiff	8
dnorm_parallel	11
dtrhpa	12
ehpa	15
etrhpa	17
hpaBinary	19
hpaML	24
hpaSelection	30
ihpa	37
ihpaDiff	39
itrhpa	43
logLik.hpaBinary	46
logLik.hpaML	46
logLik.hpaSelection	47
logLik_hpaBinary	47
logLik_hpaML	47
logLik_hpaSelection	48
mecdf	48
normalMoment	48
phpa	50
plot.hpaBinary	52
plot.hpaML	53
plot.hpaSelection	53
plot_hpaBinary	54
plot_hpaSelection	54
pnorm_parallel	55
polynomialIndex	55
predict.hpaBinary	56
predict.hpaML	57
predict.hpaSelection	58
predict_hpaBinary	59
predict_hpaML	59
predict_hpaSelection	60
print.hpaBinary	61
print.hpaML	61
print.hpaSelection	62
print.summary.hpaBinary	62
print.summary.hpaML	63
print.summary.hpaSelection	63
printPolynomial	64
print_summary_hpaBinary	64
print_summary_hpaML	65
print_summary_hpaSelection	65
summary.hpaBinary	66
summary.hpaML	66
summary.hpaSelection	67
summary_hpaBinary	67
summary_hpaML	68

<i>AIC.hpaBinary</i>	3
summary_hpaSelection	68
truncatedNormalMoment	69
Index	72

AIC.hpaBinary	<i>Calculates AIC for "hpaBinary" object</i>
---------------	--

Description

This function calculates AIC for "hpaBinary" object

Usage

```
## S3 method for class 'hpaBinary'
AIC(object, ..., k = 2)
```

Arguments

- object Object of class "hpaBinary"
- ... further arguments (currently ignored)
- k numeric, the penalty per parameter to be used; the default k = 2 is the classical AIC.

AIC.hpaML	<i>Calculates AIC for "hpaML" object</i>
-----------	--

Description

This function calculates AIC for "hpaML" object

Usage

```
## S3 method for class 'hpaML'
AIC(object, ..., k = 2)
```

Arguments

- object Object of class "hpaML"
- ... further arguments (currently ignored)
- k numeric, the penalty per parameter to be used; the default k = 2 is the classical AIC.

AIC.hpaSelection	<i>Calculates AIC for "hpaSelection" object</i>
------------------	---

Description

This function calculates AIC for "hpaSelection" object

Usage

```
## S3 method for class 'hpaSelection'  
AIC(object, ..., k = 2)
```

Arguments

object	Object of class "hpaSelection"
...	further arguments (currently ignored)
k	numeric, the penalty per parameter to be used; the default k = 2 is the classical AIC.

AIC_hpaBinary	<i>Calculates AIC for "hpaBinary" object</i>
---------------	--

Description

This function calculates AIC for "hpaBinary" object

Usage

```
AIC_hpaBinary(object, k = 2)
```

Arguments

object	Object of class "hpaBinary"
k	numeric, the penalty per parameter to be used; the default k = 2 is the classical AIC.

AIC_hpaML	<i>Calculates AIC for "hpaML" object</i>
-----------	--

Description

This function calculates AIC for "hpaML" object

Usage

```
AIC_hpaML(object, k = 2)
```

Arguments

object	Object of class "hpaML"
k	numeric, the penalty per parameter to be used; the default k = 2 is the classical AIC.

AIC_hpaSelection	<i>Calculates AIC for "hpaSelection" object</i>
------------------	---

Description

This function calculates AIC for "hpaSelection" object

Usage

```
AIC_hpaSelection(object, k = 2)
```

Arguments

object	Object of class "hpaSelection"
k	numeric, the penalty per parameter to be used; the default k = 2 is the classical AIC.

dhp

*Density function hermite polynomial approximation***Description**

This function calculates density function hermite polynomial approximation.

Usage

```
dhp(
  x = matrix(1, 1),
  pol_coefficients = numeric(0),
  pol_degrees = numeric(0),
  given_ind = logical(0),
  omit_ind = logical(0),
  mean = numeric(0),
  sd = numeric(0),
  is_parallel = FALSE,
  is_log = FALSE
)
```

Arguments

<code>x</code>	numeric matrix of density function arguments. Note that <code>x</code> rows are observations while variables are columns.
<code>pol_coefficients</code>	numeric vector of polynomial coefficients.
<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
<code>given_ind</code>	logical vector indicating wheather corresponding component is conditioned. By default it is a logical vector of FALSE values.
<code>omit_ind</code>	logical vector indicating wheather corresponding component is omitted. By default it is a logical vector of FALSE values.
<code>mean</code>	numeric vector of expected values.
<code>sd</code>	positive numeric vector of standard deviations.
<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
<code>is_log</code>	logical; if TRUE then probabilities <code>p</code> are given as $\log(p)$ or derivatives will be given respect to $\log(p)$

Details

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to

insure integration to 1) product of squared polynomial and normal distribution densities. Parameters mean and sd determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below.

Parameters mean, sd, given_ind, omit_ind should have the same length as pol_degrees parameter.

Value

This function returns density function hermite polynomial approximation at point x.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

Examples

```
## Let's approximate some three random variables joint density function
## at point (0,1, 0.2, 0.3) with hermite polynomial of (1,2,3) degrees
## which polynomial coefficients equals 1 except coefficient related
## to x1*(x^3) polynomial element which equals 2. Also suppose that normal
## density related mean vector equals (1.1, 1.2, 1.3) while standard deviations
## vector is (2.1, 2.2, 2.3).

# Prepare initial values
x <- matrix(c(0.1, 0.2, 0.3), nrow = 1)
mean <- c(1.1, 1.2, 1.3)
sd <- c(2.1, 2.2, 2.3)
pol_degrees <- c(1, 2, 3)

# Create polynomial powers and indexes correspondence matrix
pol_ind <- polynomialIndex(pol_degrees)
# Set all polynomial coefficients to 1
pol_coefficients <- rep(1, ncol(pol_ind))
pol_degrees_n <- length(pol_degrees)

# Assign coefficient 2 to the polynomial element(x1 ^ 1)*(x2 ^ 0)*(x3 ^ 2)
pol_coefficients[apply(pol_ind, 2, function(x) all(x == c(1, 0, 2)))] <- 2

# Visualize correspondence between polynomial
# elements and their coefficients
as.data.frame(rbind(pol_ind, pol_coefficients),
  row.names = c("x1 power", "x2 power", "x3 power", "coefficients"),
  optional = TRUE)
printPolynomial(pol_degrees, pol_coefficients)

# Calculate density approximation at point x
dhpa(x = x,
  pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
  mean = mean, sd = sd)

# Condition second component to be 0.5
# Substitute x second component with conditional value 0.5
```

```

x <- matrix(c(0.1, 0.5, 0.3), nrow = 1)
#Set TRUE to the second component indicating that it is conditioned
given_ind <- c(FALSE, TRUE, FALSE)

# Calculate conditional (on x2=0.5) density approximation at point x
dhpa(x = x,
     pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
     mean = mean, sd = sd,
     given_ind = given_ind)

# Consider third component marginal distribution
# conditioned on the second component 0.5 value
# Set TRUE to the first component indicating that it is omitted
omit_ind <- c(TRUE, FALSE, FALSE)

# Calculate conditional (on x2=0.5) marginal (for x3) density approximation
# at point x
dhpa(x = x,
     pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
     mean = mean, sd = sd,
     given_ind = given_ind, omit_ind = omit_ind)

```

dhpaDiff

Calculate gradient of density function hermite polynomial approximation

Description

This function calculates gradient of density function hermite polynomial approximation.

Usage

```

dhpaDiff(
  x = matrix(1, 1),
  pol_coefficients = numeric(0),
  pol_degrees = numeric(0),
  given_ind = logical(0),
  omit_ind = logical(0),
  mean = numeric(0),
  sd = numeric(0),
  type = "pol_coefficients",
  is_parallel = FALSE,
  is_log = FALSE
)

```

Arguments

x numeric matrix of density function arguments. Note that x rows are observations while variables are columns.

<code>pol_coefficients</code>	numeric vector of polynomial coefficients.
<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
<code>given_ind</code>	logical vector indicating wheather corresponding component is conditioned. By default it is a logical vector of FALSE values.
<code>omit_ind</code>	logical vector indicating wheather corresponding component is omitted. By default it is a logical vector of FALSE values.
<code>mean</code>	numeric vector of expected values.
<code>sd</code>	positive numeric vector of standard deviations.
<code>type</code>	determines the partial derivatives to be included into gradient. If <code>type="pol_coefficients"</code> then gradient will contain partial derivatives respect to polynomial coefficients listed in the same order as <code>pol_coefficients</code> . Other available types are <code>type="mean"</code> and <code>type="sd"</code> . For function <code>dhpDiff</code> it is possible to take gradient respect to the <code>x</code> points setting <code>type="x"</code> . For function <code>ihpaDiff</code> it is possible to take gradient respect to the <code>x</code> lower and upper points setting <code>type="x_lower"</code> or <code>type="upper"</code> correspondingly. In order to get full gradient please set <code>type="all"</code> .
<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
<code>is_log</code>	logical; if TRUE then probabilities <code>p</code> are given as <code>log(p)</code> or derivatives will be given respect to <code>log(p)</code>

Details

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters `mean` and `sd` determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below.

Parameters `mean`, `sd`, `given_ind`, `omit_ind` should have the same length as `pol_degrees` parameter.

If `x` has more then one row then the output will be jacobian matrix where rows are gradients.

Value

This function returns gradient of density function hermite polynomial approximation at point `x`. Gradient elements are determined by the `type` argument.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

Examples

```
## Let's approximate some three random variables joint density function
## at point (0.1, 0.2, 0.3) with hermite polynomial of (1,2,3) degrees
## which polynomial coefficients equals 1 except coefficient related to
## x1*(x^3) polynomial element which equals 2. Also suppose that normal
## density related mean vector equals (1.1, 1.2, 1.3) while standard
## deviations vector is (2.1, 2.2, 2.3). In this example let's calculate
## density approximating function's gradient respect to various parameters.

# Prepare initial values
x <- matrix(c(0.1, 0.2, 0.3), nrow = 1)
mean <- c(1.1, 1.2, 1.3)
sd <- c(2.1, 2.2, 2.3)
pol_degrees <- c(1, 2, 3)

# Create polynomial powers and indexes correspondence matrix
pol_ind <- polynomialIndex(pol_degrees)
# Set all polynomial coefficients to 1
pol_coefficients <- rep(1, ncol(pol_ind))
pol_degrees_n <- length(pol_degrees)

# Assign coefficient 2 to the polynomial element(x1 ^ 1)*(x2 ^ 0)*(x3 ^ 2)
pol_coefficients[apply(pol_ind, 2, function(x) all(x == c(1, 0, 2)))] <- 2

# Visualize correspondence between polynomial
# elements and their coefficients
as.data.frame(rbind(pol_ind, pol_coefficients),
  row.names = c("x1 power", "x2 power", "x3 power", "coefficients"),
  optional = TRUE)
printPolynomial(pol_degrees, pol_coefficients)

# Calculate density approximation gradient
# respect to polynomial coefficients at point x
dhpaDiff(x = x,
  pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
  mean = mean, sd = sd)

# Condition second component to be 0.5
# Substitute x second component with conditional value 0.5
x <- matrix(c(0.1, 0.5, 0.3), nrow = 1)
# Set TRUE to the second component indicating that it is conditioned
given_ind <- c(FALSE, TRUE, FALSE)

# Calculate conditional (on x2 = 0.5) density approximation's
# gradient respect to polynomial coefficients at point x
dhpaDiff(x = x,
  pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
  mean = mean, sd = sd,
  given_ind = given_ind)

# Consider third component marginal distribution
# conditioned on the second component 0.5 value
```

```

# Set TRUE to the first component indicating that it is omitted
omit_ind <- c(TRUE, FALSE, FALSE)

# Calculate conditional (on x2 = 0.5) marginal (for x3) density
# approximation's gradient respect to:
#   polynomial coefficients
dhpDiff(x = x,
        pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
        mean = mean, sd = sd,
        given_ind = given_ind, omit_ind = omit_ind)
#   mean
dhpDiff(x = x,
        pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
        mean = mean, sd = sd,
        given_ind = given_ind, omit_ind = omit_ind,
        type = "mean")
#   sd
dhpDiff(x = x,
        pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
        mean = mean, sd = sd,
        given_ind = given_ind, omit_ind = omit_ind,
        type = "sd")
#   x
dhpDiff(x = x,
        pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
        mean = mean, sd = sd,
        given_ind = given_ind, omit_ind = omit_ind,
        type = "x")

```

dnorm_parallel

Calculate normal pdf in parallel

Description

Calculate in parallel for each value from vector x density function of normal distribution with mean equal to mean and standard deviation equal to sd.

Usage

```
dnorm_parallel(x, mean = 0, sd = 1, is_parallel = FALSE)
```

Arguments

<code>x</code>	vector of quantiles: should be numeric vector, not just double value.
<code>mean</code>	double value.
<code>sd</code>	double positive value.
<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).

Examples

```
## Consider normal distribution with mean 3 and standard deviation 5.
## Calculate it's density function at points 2 and 3.

# Create vector of points
my_points <- c(2, 3)

# Calculate pdf at these points
# (set is_parallel = TRUE in order
# to turn on parallel computations)
dnorm_parallel(my_points, 3, 5,
               is_parallel = FALSE)
```

dtrhpa

Truncated density function hermite polynomial approximation

Description

This function calculates truncated density function hermite polynomial approximation.

Usage

```
dtrhpa(
  x = matrix(1, 1),
  tr_left = matrix(),
  tr_right = matrix(),
  pol_coefficients = numeric(0),
  pol_degrees = numeric(0),
  given_ind = logical(0),
  omit_ind = logical(0),
  mean = numeric(0),
  sd = numeric(0),
  is_parallel = FALSE,
  is_log = FALSE
)
```

Arguments

x	numeric matrix of density function arguments. Note that x rows are observations while variables are columns.
tr_left	numeric matrix of left (lower) truncation limits. Note that tr_right rows are observations while variables are columns. If tr_left or tr_right is single row matrix then the same truncation limits would be applied to all observations that are determined by the first rows of these matrices.
tr_right	numeric matrix of right (upper) truncation limits. Note that tr_right rows are observations while variables are columns. If tr_left or tr_right is single row matrix then the same truncation limits would be applied to all observations that are determined by the first rows of these matrices.

<code>pol_coefficients</code>	numeric vector of polynomial coefficients.
<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
<code>given_ind</code>	logical vector indicating wheather corresponding component is conditioned. By default it is a logical vector of FALSE values.
<code>omit_ind</code>	logical vector indicating wheather corresponding component is omitted. By default it is a logical vector of FALSE values.
<code>mean</code>	numeric vector of expected values.
<code>sd</code>	positive numeric vector of standard deviations.
<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
<code>is_log</code>	logical; if TRUE then probabilities p are given as log(p) or derivatives will be given respect to log(p)

Details

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters `mean` and `sd` determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below. Parameters `mean`, `sd`, `given_ind`, `omit_ind` should have the same length as `pol_degrees` parameter.

Value

This function returns density function hermite polynomial approximation at point `x` for truncated distribution.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

Examples

```
## Let's approximate some three random variables joint density function
## at point (0,1, 0.2, 0.3) with hermite polynomial of (1,2,3) degrees
## which polynomial coefficients equals 1 except coefficient related to
## x1*(x^3) polynomial element which equals 2. Also suppose that normal
## density related mean vector equals (1.1, 1.2, 1.3) while standard
## deviations vector is (2.1, 2.2, 2.3). Suppose that lower and upper
## truncation points are (-1.1,-1.2,-1.3) and (1.1,1.2,1.3) correspondingly.

# Prepare initial values
x <- matrix(c(0.1, 0.2, 0.3), nrow=1)
tr_left = matrix(c(-1.1,-1.2,-1.3), nrow = 1)
```

```

tr_right = matrix(c(1.1,1.2,1.3), nrow = 1)
mean <- c(1.1, 1.2, 1.3)
sd <- c(2.1, 2.2, 2.3)
pol_degrees <- c(1, 2, 3)

# Create polynomial powers and indexes correspondence matrix
pol_ind <- polynomialIndex(pol_degrees)

# Set all polynomial coefficients to 1
pol_coefficients <- rep(1, ncol(pol_ind))
pol_degrees_n <- length(pol_degrees)

# Assign coefficient 2 to the polynomial element(x1 ^ 1)*(x2 ^ 0)*(x3 ^ 2)
pol_coefficients[apply(pol_ind, 2, function(x) all(x == c(1, 0, 2)))] <- 2

# Visualize correspondence between polynomial elements and their coefficients
as.data.frame(rbind(pol_ind, pol_coefficients),
row.names = c("x1 power", "x2 power", "x3 power", "coefficients"),
optional = TRUE)
printPolynomial(pol_degrees, pol_coefficients)

# Calculate density approximation at point x
dtrhpa(x = x,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
tr_left = tr_left, tr_right = tr_right)

# Condition second component to be 0.5
# Substitute x second component with conditional value 0.5
x <- matrix(c(0.1, 0.5, 0.3), nrow = 1)
# Set TRUE to the second component indicating that it is conditioned
given_ind <- c(FALSE, TRUE, FALSE)
# Calculate conditional (on x2=0.5) density approximation at point x
dtrhpa(x = x,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
given_ind = given_ind,
tr_left = tr_left, tr_right = tr_right)

# Consider third component marginal distribution
# conditioned on the second component 0.5 value
# Set TRUE to the first component indicating that it is omitted
omit_ind <- c(TRUE, FALSE, FALSE)

# Calculate conditional (on x2=0.5) marginal (for x3) density approximation at point x
dtrhpa(x = x,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
given_ind = given_ind, omit_ind = omit_ind,
tr_left = tr_left, tr_right = tr_right)

```

ehpa

*Expected powered product hermite polynomial approximation***Description**

This function calculates expected powered product hermite polynomial approximation.

Usage

```
ehpa(
  x = matrix(1, 1),
  pol_coefficients = numeric(0),
  pol_degrees = numeric(0),
  given_ind = logical(0),
  omit_ind = logical(0),
  mean = numeric(0),
  sd = numeric(0),
  expectation_powers = numeric(0),
  is_parallel = FALSE
)
```

Arguments

<code>x</code>	non-negative numeric matrix of quantiles. Note that <code>x</code> rows are observations while variables are columns.
<code>pol_coefficients</code>	numeric vector of polynomial coefficients.
<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
<code>given_ind</code>	logical vector indicating wheather corresponding component is conditioned. By default it is a logical vector of FALSE values.
<code>omit_ind</code>	logical vector indicating wheather corresponding component is omitted. By default it is a logical vector of FALSE values.
<code>mean</code>	numeric vector of expected values.
<code>sd</code>	positive numeric vector of standard deviations.
<code>expectation_powers</code>	integer vector of random vector components powers.
<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).

Details

Expected powered product of random variables is expectation of their product given powers `expectation_powers`. Therefore in order to approximate expected value of *i*-th random vector component just set all `expectation_powers` to zero except it's *i*-th component which should be assigned 1.

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters `mean` and `sd` determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below.

Parameters `mean`, `sd`, `given_ind`, `omit_ind` should have the same length as `pol_degrees` parameter.

Value

This function returns numeric vector of expected powered product hermite polynomial approximations.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

Examples

```
## Let's approximate some three random variables powered product
## expectation for powers (3,2,1) with hermite polynomial of (1,2,3)
## degrees which polynomial coefficients equals 1 except coefficient
## related to x1*(x^3) polynomial element which equals 2.
## Also suppose that normal density related mean vector equals
## (1.1, 1.2, 1.3) while standard deviations vector is (2.1, 2.2, 2.3).

# Prepare initial values
expectation_powers = c(3,2,1)
mean <- c(1.1, 1.2, 1.3)
sd <- c(2.1, 2.2, 2.3)
pol_degrees <- c(1, 2, 3)

# Create polynomial powers and indexes correspondence matrix
pol_ind <- polynomialIndex(pol_degrees)

# Set all polynomial coefficients to 1
pol_coefficients <- rep(1, ncol(pol_ind))
pol_degrees_n <- length(pol_degrees)

# Assign coefficient 2 to the polynomial element (x1 ^ 1)*(x2 ^ 0)*(x3 ^ 2)
pol_coefficients[which(colSums(pol_ind == c(1, 0, 2)) == pol_degrees_n)] <- 2

# Visualize correspondence between polynomial elements and their coefficients
as.data.frame(rbind(pol_ind, pol_coefficients),
  row.names = c("x1 power", "x2 power", "x3 power", "coefficients"),
  optional = TRUE)
printPolynomial(pol_degrees, pol_coefficients)

# Calculate expected powered product approximation
ehpa(pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
```

```

mean = mean, sd = sd, expectation_powers = expectation_powers)

# Condition second component to be 0.5
# Substitute x second component with conditional value 0.5
x <- matrix(c(NA, 0.5, NA), nrow = 1)
#Set TRUE to the second component indicating that it is conditioned
given_ind <- c(FALSE, TRUE, FALSE)

# Calculate conditional(on x2 = 0.5) expected powered product approximation
ehpa(x = x,
    pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
    mean = mean, sd = sd, expectation_powers = expectation_powers,
    given_ind = given_ind)

# Consider third component marginal distribution
# conditioned on the second component 0.5 value
# Set TRUE to the first component indicating that it is omitted
omit_ind <- c(TRUE, FALSE, FALSE)

# Calculate conditional (on x2=0.5) marginal (for x3) expected powered
# product approximation at points x_lower and x_upper
ehpa(x = x,
    pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
    mean = mean, sd = sd, expectation_powers = expectation_powers,
    given_ind = given_ind, omit_ind = omit_ind)

```

etrhpa

Expected powered product hermite polynomial approximation for truncated distribution

Description

This function calculates expected powered product hermite polynomial approximation for truncated distribution.

Usage

```

etrhpa(
  tr_left = matrix(1, 1),
  tr_right = matrix(1, 1),
  pol_coefficients = numeric(0),
  pol_degrees = numeric(0),
  mean = numeric(0),
  sd = numeric(0),
  expectation_powers = numeric(0),
  is_parallel = FALSE
)

```

Arguments

<code>tr_left</code>	numeric matrix of left (lower) truncation limits. Note that <code>tr_right</code> rows are observations while variables are columns. If <code>tr_left</code> or <code>tr_right</code> is single row matrix then the same truncation limits would be applied to all observations that are determined by the first rows of these matrices.
<code>tr_right</code>	numeric matrix of right (upper) truncation limits. Note that <code>tr_right</code> rows are observations while variables are columns. If <code>tr_left</code> or <code>tr_right</code> is single row matrix then the same truncation limits would be applied to all observations that are determined by the first rows of these matrices.
<code>pol_coefficients</code>	numeric vector of polynomial coefficients.
<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
<code>mean</code>	numeric vector of expected values.
<code>sd</code>	positive numeric vector of standard deviations.
<code>expectation_powers</code>	integer vector of random vector components powers.
<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).

Details

Expected powered product of random variables is expectation of their product given powers `expectation_powers`. Therefore in order to approximate expected value of *i*-th random vector component just set all `expectation_powers` to zero except it's *i*-th component which should be assigned 1.

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters `mean` and `sd` determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below.

Parameters `mean`, `sd`, `given_ind`, `omit_ind` should have the same length as `pol_degrees` parameter.

Value

This function returns numeric vector of expected powered product hermite polynomial approximations for truncated distribution.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

Examples

```
## Let's approximate some three truncated random variables powered
## product expectation for powers (3,2,1) with hermite polynomial of
## (1,2,3) degrees which polynomial coefficients equals 1 except coefficient
## related to  $x_1(x^3)$  polynomial element which equals 2. Also suppose that
## normal density related mean vector equals (1.1, 1.2, 1.3) while standard
## deviations vector is (2.1, 2.2, 2.3). Suppose that lower and upper
## truncation points are (-1.1,-1.2,-1.3) and (1.1,1.2,1.3) correspondingly.

# Prepare initial values
expectation_powers = c(3,2,1)
tr_left = matrix(c(-1.1,-1.2,-1.3), nrow = 1)
tr_right = matrix(c(1.1,1.2,1.3), nrow = 1)
mean <- c(1.1, 1.2, 1.3)
sd <- c(2.1, 2.2, 2.3)
pol_degrees <- c(1, 2, 3)

# Create polynomial powers and indexes correspondence matrix
pol_ind <- polynomialIndex(pol_degrees)
# Set all polynomial coefficients to 1
pol_coefficients <- rep(1, ncol(pol_ind))
pol_degrees_n <- length(pol_degrees)

# Assign coefficient 2 to the polynomial element( $x_1^1$ )*( $x_2^0$ )*( $x_3^2$ )
pol_coefficients[apply(pol_ind, 2, function(x) all(x == c(1, 0, 2)))] <- 2

# Visualize correspondence between polynomial elements and their coefficients
as.data.frame(rbind(pol_ind, pol_coefficients),
  row.names = c("x1 power", "x2 power", "x3 power", "coefficients"),
  optional = TRUE)
printPolynomial(pol_degrees, pol_coefficients)

# Calculate expected powered product approximation for truncated distribution
ethrpa(pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
  mean = mean, sd = sd, expectation_powers = expectation_powers,
  tr_left = tr_left, tr_right = tr_right)
```

hpaBinary

Perform semi-nonparametric binary choice model estimation

Description

This function performs semi-nonparametric single index binary choice model estimation via hermite polynomial densities approximation.

Usage

```
hpaBinary(
  formula,
```

```

data,
K = 1L,
z_mean_fixed = NA_real_,
z_sd_fixed = NA_real_,
z_constant_fixed = 0,
is_z_coef_first_fixed = TRUE,
is_x0_probit = TRUE,
is_sequence = FALSE,
x0 = numeric(0),
cov_type = "sandwich",
boot_iter = 100L,
is_parallel = FALSE,
opt_type = "optim",
opt_control = NULL
)

```

Arguments

formula	an object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted. All variables in formula should be numeric vectors of the same length.
data	data frame containing the variables in the model.
K	non-negative integer representing polynomial degree.
z_mean_fixed	numeric value for binary choice equation random error density mean parameter. Set it to NA (default) if this parameter should be estimated rather than fixed.
z_sd_fixed	numeric value for binary choice equation random error density sd parameter. Set it to NA (default) if this parameter should be estimated rather than fixed.
z_constant_fixed	numeric value for binary choice equation constant parameter. Set it to NA (default) if this parameter should be estimated rather than fixed.
is_z_coef_first_fixed	bool value indicating whether binary equation first independent variable coefficient should be fixed (TRUE) or estimated (FALSE).
is_x0_probit	logical; if TRUE (default) then initial points for optimization routine will be obtained by probit model estimated via glm function.
is_sequence	if TRUE then function calculates models with polynomial degrees from 0 to K each time using initial values obtained from the previous step. In this case function will return the list of models where i-th list element correspond to model calculated under K=(i-1).
x0	numeric vector of optimization routine initial values. Note that x0=c(pol_coefficients[-1],mean,sd,
cov_type	string value determining the type of covariance matrix to be returned and used for summary. If cov_type = "hessian" then negative inverse of Hessian matrix will be applied. If cov_type = "gop" then inverse of Jacobian outer products will be used. If cov_type = "sandwich" (default) then sandwich covariance matrix estimator will be applied. If cov_type = "bootstrap" then bootstrap with boot_iter iterations will be used. If cov_type = "hessianFD" or

	cov_type = "sandwichFD" then accurate but computationally demanding central difference Hessian approximation will be calculated for the inverse Hessian and sandwich estimators correspondingly. Central differences are computed via analytically provided gradient. This Hessian matrix estimation approach seems to be less accurate than BFGS approximation if polynomial order is high (usually greater than 5).
boot_iter	the number of bootstrap iterations for cov_type = "bootstrap" covariance matrix estimator type.
is_parallel	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
opt_type	string value determining the type of the optimization routine to be applied. The default is "optim" meaning that BFGS method from the optim function will be applied. If opt_type = "GA" then ga function will be additionally applied.
opt_control	a list containing arguments to be passed to the optimization routine depending on opt_type argument value. Please see details to get additional information.

Details

Semi-nonparametric (SNP) approach has been implemented via densities hermite polynomial approximation

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters `mean` and `sd` determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below.

Parameters `mean`, `sd`, `given_ind`, `omit_ind` should have the same length as `pol_degrees` parameter.

The first polynomial coefficient (zero powers) set to 1 for identification reasons.

Note that if `is_z_coef_first_fixed` value is TRUE then the coefficient for the first independent variable in formula will be fixed to 1.

If `is_x0_probit` = TRUE then parameter `sd` will be scale adjusted in order to provide better initial point for optimization routine. Please, extract `sd` adjusted value from this function's output list.

All variables mentioned in formula should be numeric vectors.

The function calculates standard errors via sandwich estimator and significance levels are reported taking into account quasi maximum likelihood estimator (QMLE) asymptotic normality. If one wants to switch from QMLE to semi-nonparametric estimator (SNPE) during hypothesis testing then covariance matrix should be estimated again using bootstrap.

This function maximizes (quasi) log-likelihood function via [optim](#) function setting its method argument to "BFGS". If `opt_type` = "GA" then genetic algorithm from [ga](#) function will be additionally (after [optim](#) putting its solution (`par`) to suggestions matrix) applied in order to perform global optimization. Note that global optimization takes much more time (usually minutes but sometimes hours or even days). The number of iterations and population size of the genetic algorithm will grow linearly along with the number of estimated parameters. If it seems that global

maximum has not been found than it is possible to continue the search restarting the function setting it's input argument `x0` to `x1` output value. Note that if `cov_type = "bootstrap"` then `ga` function will not be used for bootstrap iterations since it may be extremely time consuming.

If `opt_type = "GA"` then `opt_control` should be the list containing the values to be passed to `ga` function. It is possible to pass arguments `lower`, `upper`, `popSize`, `pcrossover`, `pmutation`, `elitism`, `maxiter`, `suggestions`, `optim`, `optimArgs`, `seed` and `monitor`. Note that it is possible to set population, selection, crossover and mutation arguments changing `ga` default parameters via `gaControl` function. These arguments information reported in `ga`. In order to provide manual values for lower and upper bounds please follow parameters ordering mentioned above for the `x0` argument. If these bonds are not provided manually then they (except those related to the polynomial coefficients) will depend on the estimates obtained by local optimization via `optim` function (this estimates will be in the middle between lower and upper). Specifically for each sd parameter lower (upper) bound is 5 times lower (higher) then this parameter `optim` estimate. For each mean and regression coefficient parameter it's lower and upper bounds deviate from corresponding `optim` estimate by two absolute value of this estimate. Finally, lower and upper bounds for each polynomial coefficient are `-10` and `10` correspondingly (do not depend on their `optim` estimates).

The following arguments are differ from their defaults in `ga`:

- `pmutation = 0.2`,
- `optim = TRUE`,
- `optimArgs = list("method" = "Nelder-Mead", "poptim" = 0.2, "pressel" = 0.5)`,
- `seed = 8`,
- `elitism = 2 + round(popSize * 0.1)`.

Let's denote by `n_reg` the number of regressors included to the formula. The arguments `popSize` and `maxiter` of `ga` function have been set proportional to the number of estimated polynomial coefficients and independent variables:

- `popSize = 10 + 5 * (K + 1) + 2 * n_reg`
- `maxiter = 50 * (1 + K) + 10 * n_reg`

Value

This function returns an object of class "hpaBinary".

An object of class "hpaBinary" is a list containing the following components:

- `optim` - `optim` function output. If `opt_type = "GA"` then it is the list containing `optim` and `ga` functions outputs.
- `x1` - numeric vector of distribution parameters estimates.
- `mean` - mean (μ) parameter of density function estimate.
- `sd` - sd (σ) parameter of density function estimate.
- `pol_coefficients` - polynomial coefficients estimates.
- `pol_degrees` - the same as `K` input parameter.
- `coefficients` - regression (single index) coefficients estimates.

- `cov_mat` - covariance matrix estimate.
- `marginal_effects` - marginal effects matrix where columns are variables and rows are observations.
- `results` - numeric matrix representing estimation results.
- `log-likelihood` - value of Log-Likelihood function.
- `AIC` - AIC value.
- `errors_exp` - random error expectation estimate.
- `errors_var` - random error variance estimate.
- `dataframe` - dataframe containing variables mentioned in formula without NA values.
- `model_Lists` - lists containing information about fixed parameters and parameters indexes in `x1`.
- `n_obs` - number of observations.
- `z_latent` - latent variable (single index) estimates.
- `z_prob` - probabilities of positive outcome (i.e. 1) estimates.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

See Also

[summary.hpaBinary](#), [predict.hpaBinary](#), [plot.hpaBinary](#), [AIC.hpaBinary](#), [logLik.hpaBinary](#)

Examples

```
## Estimate survival probability on Titanic

library("titanic")

# Prepare data set converting
# all variables to numeric vectors
h <- data.frame("male" = as.numeric(titanic_train$Sex == "male"))
h$class_1 <- as.numeric(titanic_train$Pclass == 1)
h$class_2 <- as.numeric(titanic_train$Pclass == 2)
h$class_3 <- as.numeric(titanic_train$Pclass == 3)
h$sibl <- titanic_train$SibSp
h$survived <- titanic_train$Survived
h$age <- titanic_train$Age
h$parch <- titanic_train$Parch
h$fare <- titanic_train$Fare

# Estimate model parameters
model_hpa_1 <- hpaBinary(survived ~class_1 + class_2 +
male + age + sibl + parch + fare,
K = 3, data = h)
#get summary
summary(model_hpa_1)
```

```

# Get predicted probabilities
pred_hpa_1 <- predict(model_hpa_1)

# Calculate number of correct predictions
hpa_1_correct_0 <- sum((pred_hpa_1 < 0.5) & (model_hpa_1$dataframe$survived == 0))
hpa_1_correct_1 <- sum((pred_hpa_1 >= 0.5) & (model_hpa_1$dataframe$survived == 1))
hpa_1_correct <- hpa_1_correct_1 + hpa_1_correct_0

# Plot random errors density approximation
plot(model_hpa_1)

## Estimate parameters on data simulated from student distribution

library("mvtnorm")
set.seed(123)

# Simulate independent variables from normal distribution
n <- 5000
X <- rmvnorm(n=n, mean = c(0,0),
sigma = matrix(c(1,0.5,0.5,1), ncol=2))

# Simulate random errors from student distribution
epsilon <- rt(n, 5) * (3 / sqrt(5))

# Calculate latent and observable variables values
z_star <- 1 + X[, 1] + X[, 2] + epsilon
z <- as.numeric((z_star > 0))

# Store the results into dataframe
h <- as.data.frame(cbind(z,X))
names(h) <- c("z", "x1", "x2")

# Estimate model parameters
model <- hpaBinary(formula = z ~ x1 + x2, data=h, K = 3)
summary(model)

# Get predicted probabilities of 1 values
predict(model)

# Plot density function approximation
plot(model)

```

Description

This function performs semi-nonparametric maximum likelihood estimation via hermite polynomial densities approximation.

Usage

```
hpaML(
  x,
  pol_degrees = numeric(0),
  tr_left = numeric(0),
  tr_right = numeric(0),
  given_ind = logical(0),
  omit_ind = logical(0),
  x0 = numeric(0),
  cov_type = "sandwich",
  boot_iter = 100L,
  is_parallel = FALSE,
  opt_type = "optim",
  opt_control = NULL
)
```

Arguments

<code>x</code>	numeric matrix which rows are realizations of independent identically distributed random vectors while columns correspond to variables.
<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
<code>tr_left</code>	numeric matrix of left (lower) truncation limits. Note that <code>tr_right</code> rows are observations while variables are columns. If <code>tr_left</code> or <code>tr_right</code> is single row matrix then the same truncation limits would be applied to all observations that are determined by the first rows of these matrices.
<code>tr_right</code>	numeric matrix of right (upper) truncation limits. Note that <code>tr_right</code> rows are observations while variables are columns. If <code>tr_left</code> or <code>tr_right</code> is single row matrix then the same truncation limits would be applied to all observations that are determined by the first rows of these matrices.
<code>given_ind</code>	logical vector indicating wheather corresponding component is conditioned. By default it is a logical vector of FALSE values.
<code>omit_ind</code>	logical vector indicating wheather corresponding component is omitted. By default it is a logical vector of FALSE values.
<code>x0</code>	numeric vector of optimization routine initial values. Note that <code>x0=c(pol_coefficients[-1],mean,sd)</code> For <code>pol_coefficients</code> , <code>mean</code> and <code>sd</code> documentation see dhp function.
<code>cov_type</code>	string value determining the type of covariance matrix to be returned and used for summary. If <code>cov_type = "hessian"</code> then negative inverse of Hessian matrix will be applied. If <code>cov_type = "gop"</code> then inverse of Jacobian outer products will be used. If <code>cov_type = "sandwich"</code> (default) then sandwich covariance matrix estimator will be applied. If <code>cov_type = "bootstrap"</code> then bootstrap with <code>boot_iter</code> iterations will be used. If <code>cov_type = "hessianFD"</code> or

	cov_type = "sandwichFD" then accurate but computationally demanding central difference Hessian approximation will be calculated for the inverse Hessian and sandwich estimators correspondingly. Central differences are computed via analytically provided gradient. This Hessian matrix estimation approach seems to be less accurate than BFGS approximation if polynomial order is high (usually greater than 5).
boot_iter	the number of bootstrap iterations for cov_type = "bootstrap" covariance matrix estimator type.
is_parallel	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
opt_type	string value determining the type of the optimization routine to be applied. The default is "optim" meaning that BFGS method from the optim function will be applied. If opt_type = "GA" then ga function will be additionally applied.
opt_control	a list containing arguments to be passed to the optimization routine depending on opt_type argument value. Please see details to get additional information.

Details

Semi-nonparametric (SNP) approach has been implemented via densities hermite polynomial approximation

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters mean and sd determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below.

Parameters mean, sd, given_ind, omit_ind should have the same length as `pol_degrees` parameter.

The first polynomial coefficient (zero powers) set to 1 for identification reasons.

The function calculates standard errors via sandwich estimator and significance levels are reported taking into account quasi maximum likelihood estimator (QMLE) asymptotic normality. If one wants to switch from QMLE to semi-nonparametric estimator (SNPE) during hypothesis testing then covariance matrix should be estimated again using bootstrap.

This function maximizes (quasi) log-likelihood function via [optim](#) function setting its method argument to "BFGS". If opt_type = "GA" then genetic algorithm from [ga](#) function will be additionally (after [optim](#) putting its solution (par) to suggestions matrix) applied in order to perform global optimization. Note that global optimization takes much more time (usually minutes but sometimes hours or even days). The number of iterations and population size of the genetic algorithm will grow linearly along with the number of estimated parameters. If it seems that global maximum has not been found then it is possible to continue the search restarting the function setting its input argument `x0` to `x1` output value. Note that if cov_type = "bootstrap" then [ga](#) function will not be used for bootstrap iterations since it may be extremely time consuming.

If opt_type = "GA" then opt_control should be the list containing the values to be passed to [ga](#) function. It is possible to pass arguments lower, upper, popSize, pcrossover, pmutation, elitism, maxiter, suggestions, optim, optimArgs, seed and monitor. Note that it is possible

to set population, selection, crossover and mutation arguments changing `ga` default parameters via `gaControl` function. These arguments information reported in `ga`. In order to provide manual values for lower and upper bounds please follow parameters ordering mentioned above for the `x0` argument. If these bonds are not provided manually then they (except those related to the polynomial coefficients) will depend on the estimates obtained by local optimization via `optim` function (this estimates will be in the middle between lower and upper). Specifically for each `sd` parameter lower (upper) bound is 5 times lower (higher) then this parameter `optim` estimate. For each mean and regression coefficient parameter it's lower and upper bounds deviate from corresponding `optim` estimate by two absolute value of this estimate. Finally, lower and upper bounds for each polynomial coefficient are -10 and 10 correspondingly (do not depend on their `optim` estimates).

The following arguments are differ from their defaults in `ga`:

- `pmutation = 0.2`,
- `optim = TRUE`,
- `optimArgs = list("method" = "Nelder-Mead", "poptim" = 0.2, "pressel" = 0.5)`,
- `seed = 8`,
- `elitism = 2 + round(popSize * 0.1)`.

The arguments `popSize` and `maxiter` of `ga` function have been set proportional to the number of estimated polynomial coefficients

- `popSize = 10 + (prod(pol_degrees + 1) - 1) * 2`.
- `maxiter = 50 * (prod(pol_degrees + 1))`

Value

This function returns an object of class "hpaML".

An object of class "hpaML" is a list containing the following components:

- `optim` - `optim` function output. If `opt_type = "GA"` then it is the list containing `optim` and `ga` functions outputs.
- `x1` - numeric vector of distribution parameters estimates.
- `mean` - density function mean vector estimate.
- `sd` - density function sd vector estimate.
- `pol_coefficients` - polynomial coefficients estimates.
- `tr_left` - the same as `tr_left` input parameter.
- `tr_right` - the same as `tr_right` input parameter.
- `omit_ind` - the same as `omit_ind` input parameter.
- `given_ind` - the same as `given_ind` input parameter.
- `cov_mat` - covariance matrix estimate.
- `results` - numeric matrix representing estimation results.
- `log-likelihood` - value of Log-Likelihood function.
- `AIC` - AIC value.

- data - the same as x input parameter but without NA observations.
- n_obs - number of observations.
- bootstrap - list where bootstrap estimation results are stored.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

See Also

[summary.hpaML](#), [predict.hpaML](#), [AIC.hpaML](#), [logLik.hpaML](#)

Examples

```
## Approximate student (t) distribution

# Set seed for reproducibility
set.seed(123)

# Simulate 5000 realizations of student distribution with 5 degrees of freedom
n <- 5000
df <- 5
x <- matrix(rt(n, df), ncol = 1)
pol_degrees <- c(4)

# Apply pseudo maximum likelihood routine
ml_result <- hpa::hpaML(x = x, pol_degrees = pol_degrees)
summary(ml_result)

# Get predicted probabilities (density values) approximations
predict(ml_result)

# Plot density approximation
plot(ml_result)

## Approximate chi-squared distribution

# Set seed for reproducibility
set.seed(123)

# Simulate 5000 realizations of chi-squared distribution with 5 degrees of freedom

n <- 5000
df <- 5
x <- matrix(rchisq(n, df), ncol = 1)
pol_degrees <- c(5)

# Apply pseudo maximum likelihood routine
ml_result <- hpaML(x = x, pol_degrees = as.vector(pol_degrees),
tr_left = 0)
summary(ml_result)
```

```

# Get predicted probabilities (density values) approximations
predict(ml_result)

# Plot density approximation
plot(ml_result)

## Approximate multivariate student (t) distribution
## Note that calculations may take up to a minute

# Set seed for reproducibility
set.seed(123)

# Simulate 5000 realizations of three dimensional student distribution with 5 degrees of freedom
library("mvtnorm")
cov_mat <- matrix(c(1, 0.5, -0.5, 0.5, 1, 0.5, -0.5, 0.5, 1), ncol = 3)
x <- rmvt(n = 5000, sigma = cov_mat, df = 5)

# Estimate approximating joint distribution parameters
ml_result <- hpaML(x = x, pol_degrees = c(1, 1, 1))

# Get summary
summary(ml_result)

# Get predicted values for joint density function
predict(ml_result)

# Plot density approximation for the
# second random variable
plot(ml_result, ind = 2)

# Plot density approximation for the
# second random variable conditioning
# on x1 = 1
plot(ml_result, ind = 2, given = c(1, NA, NA))

## Approximate student (t) distribution and plot densities approximated
## under different hermite polynomial degrees against
## true density (of student distribution)

# Simulate 5000 realizations of t-distribution with 5 degrees of freedom
n <- 5000
df <- 5
x <- matrix(rt(n, df), ncol=1)

# Apply pseudo maximum likelihood routine
# Create matrix of lists where i-th element contains hpaML results for K=i
ml_result <- matrix(list(), 4, 1)
for(i in 1:4)
{
  ml_result[[i]] <- hpa::hpaML(x = x, pol_degrees = i)
}

```

```

# Generate test values
test_values <- seq(qt(0.001, df), qt(0.999, df), 0.001)
n0 <- length(test_values)

# t-distribution density function at test values points
true_pred <- dt(test_values, df)

# Create matrix of lists where i-th element contains densities predictions for K=i
PGN_pred <- matrix(list(), 4, 1)
for(i in 1:4)
{
  PGN_pred[[i]] <- predict(object = ml_result[[i]],
                          newdata = matrix(test_values, ncol=1))
}
# Plot the result
library("ggplot2")

# prepare the data
h <- data.frame("values" = rep(test_values,5),
                "predictions" = c(PGN_pred[[1]],PGN_pred[[2]],
                                  PGN_pred[[3]],PGN_pred[[4]],
                                  true_pred),
                "Density" = c(
                  rep("K=1",n0), rep("K=2",n0),
                  rep("K=3",n0), rep("K=4",n0),
                  rep("t-distribution",n0))
                )

# build the plot
ggplot(h, aes(values, predictions)) + geom_point(aes(color = Density)) +
  theme_minimal() + theme(legend.position = "top", text = element_text(size=26),
                          legend.title=element_text(size=20), legend.text=element_text(size=28)) +
  guides(colour = guide_legend(override.aes = list(size=10))
  )

# Get informative estimates summary for K=4
summary(ml_result[[4]])

```

hpaSelection

Perform semi-nonparametric selection model estimation

Description

This function performs semi-nonparametric selection model estimation via hermite polynomial densities approximation.

Usage

```
hpaSelection(
  selection,
  outcome,
  data,
  z_K = 1L,
  y_K = 1L,
  pol_elements = 3L,
  is_Newey = FALSE,
  x0 = numeric(0),
  is_Newey_loocv = FALSE,
  cov_type = "sandwich",
  boot_iter = 100L,
  is_parallel = FALSE,
  opt_type = "optim",
  opt_control = NULL
)
```

Arguments

selection	an object of class "formula" (or one that can be coerced to that class): a symbolic description of the selection equation form. All variables in selection should be numeric vectors of the same length.
outcome	an object of class "formula" (or one that can be coerced to that class): a symbolic description of the outcome equation form. All variables in outcome should be numeric vectors of the same length.
data	data frame containing the variables in the model.
z_K	non-negative integer representing polynomial degree related to selection equation.
y_K	non-negative integer representing polynomial degree related to outcome equation.
pol_elements	number of conditional expectation approximating terms for Newey's method. If is_Newey_loocv is TRUE then determines maximum number of these terms during leave-one-out cross-validation.
is_Newey	logical; if TRUE then returns only Newey's method estimation results (default value is FALSE).
x0	numeric vector of optimization routine initial values. Note that x0=c(pol_coefficients[-1],mean,sd,
is_Newey_loocv	logical; if TRUE then number of conditional expectation approximating terms for Newey's method will be selected based on leave-one-out cross-validation criteria iterating through 0 to pol_elements number of these terms.
cov_type	string value determining the type of covariance matrix to be returned and used for summary. If cov_type = "hessian" then negative inverse of Hessian matrix will be applied. If cov_type = "gop" then inverse of Jacobian outer products will be used. If cov_type = "sandwich" (default) then sandwich covariance matrix estimator will be applied. If cov_type = "bootstrap" then bootstrap with boot_iter iterations will be used. If cov_type = "hessianFD" or

	cov_type = "sandwichFD" then accurate but computationally demanding central difference Hessian approximation will be calculated for the inverse Hessian and sandwich estimators correspondingly. Central differences are computed via analytically provided gradient. This Hessian matrix estimation approach seems to be less accurate than BFGS approximation if polynomial order is high (usually greater than 5).
boot_iter	the number of bootstrap iterations for cov_type = "bootstrap" covariance matrix estimator type.
is_parallel	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
opt_type	string value determining the type of the optimization routine to be applied. The default is "optim" meaning that BFGS method from the optim function will be applied. If opt_type = "GA" then ga function will be additionally applied.
opt_control	a list containing arguments to be passed to the optimization routine depending on opt_type argument value. Please see details to get additional information.

Details

Semi-nonparametric (SNP) approach has been implemented via densities hermite polynomial approximation

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree pol_degree. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters mean and sd determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below.

Parameters mean, sd, given_ind, omit_ind should have the same length as pol_degrees parameter.

The first polynomial coefficient (zero powers) set to 1 for identification reasons.

Note that coefficient for the first independent variable in selection will be fixed to 1.

All variables mentioned in selection and outcome should be numeric vectors.

The function calculates standard errors via sandwich estimator and significance levels are reported taking into account quasi maximum likelihood estimator (QMLE) asymptotic normality. If one wants to switch from QMLE to semi-nonparametric estimator (SNPE) during hypothesis testing then covariance matrix should be estimated again using bootstrap.

Initial values for optimization routine are obtained by Newey's method (see the reference below).

Note that selection equation dependent variables should have exactly two levels (0 and 1) where "0" states for the selection results which leads to unobservable values of dependent variable in outcome equation.

This function maximizes (quasi) log-likelihood function via [optim](#) function setting its method argument to "BFGS". If opt_type = "GA" then genetic algorithm from [ga](#) function will be additionally (after [optim](#) putting its solution (par) to suggestions matrix) applied in order to perform global optimization. Note that global optimization takes much more time (usually minutes but

sometimes hours or even days). The number of iterations and population size of the genetic algorithm will grow linearly along with the number of estimated parameters. If it seems that global maximum has not been found then it is possible to continue the search restarting the function setting it's input argument `x0` to `x1` output value. Note that if `cov_type = "bootstrap"` then `ga` function will not be used for bootstrap iterations since it may be extremely time consuming.

If `opt_type = "GA"` then `opt_control` should be the list containing the values to be passed to `ga` function. It is possible to pass arguments `lower`, `upper`, `popSize`, `pcrossover`, `pmutation`, `elitism`, `maxiter`, `suggestions`, `optim`, `optimArgs`, `seed` and `monitor`. Note that it is possible to set population, selection, crossover and mutation arguments changing `ga` default parameters via `gaControl` function. These arguments information reported in `ga`. In order to provide manual values for lower and upper bounds please follow parameters ordering mentioned above for the `x0` argument. If these bonds are not provided manually then they (except those related to the polynomial coefficients) will depend on the estimates obtained by local optimization via `optim` function (this estimates will be in the middle between lower and upper). Specifically for each sd parameter lower (upper) bound is 5 times lower (higher) then this parameter `optim` estimate. For each mean and regression coefficient parameter it's lower and upper bounds deviate from corresponding `optim` estimate by two absolute value of this estimate. Finally, lower and upper bounds for each polynomial coefficient are `-10` and `10` correspondingly (do not depend on their `optim` estimates).

The following arguments differ from their defaults in `ga`:

- `pmutation = 0.2`,
- `optim = TRUE`,
- `optimArgs = list("method" = "Nelder-Mead", "poptim" = 0.2, "pressel" = 0.5)`,
- `seed = 8`,
- `elitism = 2 + round(popSize * 0.1)`.

Let's denote by `n_reg` the number of regressors included to the selection and outcome formulas. The arguments `popSize` and `maxiter` of `ga` function have been set proportional to the number of estimated polynomial coefficients and independent variables:

- `popSize = 10 + 5 * (z_K + 1) * (y_K + 1) + 2 * n_reg`
- `maxiter = 50 * (z_K + 1) * (y_K + 1) + 10 * n_reg`

Value

This function returns an object of class "hpaSelection".

An object of class "hpaSelection" is a list containing the following components:

- `optim` - `optim` function output. If `opt_type = "GA"` then it is the list containing `optim` and `ga` functions outputs.
- `x1` - numeric vector of distribution parameters estimates.
- `Newey` - list containing information concerning Newey's method estimation results.
- `z_mean` - estimate of the hermite polynomial mean parameter related to selection equation random error marginal distribution.

- `y_mean` - estimate of the hermite polynomial mean parameter related to outcome equation random error marginal distribution.
- `z_sd` - estimate of sd parameter related to selection equation random error marginal distribution.
- `y_sd` - estimate of the hermite polynomial sd parameter related to outcome equation random error marginal distribution.
- `pol_coefficients` - polynomial coefficients estimates.
- `pol_degrees` - numeric vector which first element is `z_K` and the second is `y_K`.
- `z_coef` - selection equation regression coefficients estimates.
- `y_coef` - outcome equation regression coefficients estimates.
- `cov_mat` - covariance matrix estimate.
- `results` - numeric matrix representing estimation results.
- `log-likelihood` - value of Log-Likelihood function.
- `re_moments` - list which contains information about random errors expectations, variances and correlation.
- `data_List` - list containing model variables and their partition according to outcome and selection equations.
- `n_obs` - number of observations.
- `ind_List` - list which contains information about parameters indexes in `x1`.
- `selection_formula` - the same as selection input parameter.
- `outcome_formula` - the same as outcome input parameter.

Abovementioned list `Newey` has class "hpaNewey" and contains the following components:

- `y_coef` - regression coefficients estimates (except constant term which is part of conditional expectation approximating polynomial).
- `z_coef` - regression coefficients estimates related to selection equation.
- `constant_biased` - biased estimate of constant term.
- `inv_mills` - inverse mills ratios estimates and their powers (including constant).
- `inv_mills_coef` - coefficients related to `inv_mills`.
- `pol_elements` - the same as `pol_elements` input parameter. However if `is_Newey_loocv` is TRUE then it will equal to the number of conditional expectation approximating terms for Newey's method which minimize leave-one-out cross-validation criteria.
- `outcome_exp_cond` - dependend variable conditional expectation estimates.
- `selection_exp` - selection equation random error expectation estimate.
- `selection_var` - selection equation random error variance estimate.
- `hpaBinaryModel` - object of class "hpaBinary" which contains selection equation estimation results.

Abovementioned list `re_moments` contains the following components:

- `selection_exp` - selection equation random errors expectation estimate.

- selection_var - selection equation random errors variance estimate.
- outcome_exp - outcome equation random errors expectation estimate.
- outcome_var - outcome equation random errors variance estimate.
- errors_covariance - outcome and selection equation random errors covariance estimate.
- rho - outcome and selection equation random errors correlation estimate.
- rho_std - outcome and selection equation random errors correlation estimator standard error estimate.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>
 W. K. Newey (2009) <<https://doi.org/10.1111/j.1368-423X.2008.00263.x>>
 Mroz T. A. (1987) <doi:10.2307/1911029>

See Also

[summary.hpaSelection](#), [predict.hpaSelection](#), [plot.hpaSelection](#), [AIC.hpaSelection](#), [logLik.hpaSelection](#)

Examples

```
## Let's estimate wage equation accounting for non-random selection.
## See the reference to Mroz TA (1987) to get additional details about
## the data this examples use

# Prepare data
library("sampleSelection")
data("Mroz87")
h = data.frame("kids" = as.numeric(Mroz87$kids5 + Mroz87$kids618 > 0),
  "age" = as.numeric(Mroz87$age),
  "faminc" = as.numeric(Mroz87$faminc),
  "educ" = as.numeric(Mroz87$educ),
  "exper" = as.numeric(Mroz87$exper),
  "city" = as.numeric(Mroz87$city),
  "wage" = as.numeric(Mroz87$wage),
  "lfp" = as.numeric(Mroz87$lfp))

# Estimate model parameters
model <- hpaSelection(selection = lfp ~ educ + age + I(age ^ 2) +
  kids + log(faminc),
  outcome = log(wage) ~ exper + I(exper ^ 2) +
  educ + city,
  z_K = 2, y_K = 3, data = h,
  pol_elements = 3, is_Newey_loocv = TRUE)

summary(model)

# Plot outcome equation random errors density
plot(model, is_outcome = TRUE)
# Plot selection equation random errors density
```

```

plot(model, is_outcome = FALSE)

## Estimate semi-nonparametric sample selection model
## parameters on simulated data given chi-squared random errors

set.seed(100)
library("mvtnorm")

# Sample size

n <- 1000

# Simulate independent variables
X_rho <- 0.5
X_sigma <- matrix(c(1,X_rho,X_rho,X_rho,1,X_rho,X_rho,X_rho,1), ncol=3)
X <- rmvnorm(n=n, mean = c(0,0,0),
             sigma = X_sigma)

# Simulate random errors
epsilon <- matrix(0, n, 2)
epsilon_z_y <- rchisq(n, 5)
epsilon[, 1] <- (rchisq(n, 5) + epsilon_z_y) * (sqrt(3/20)) - 3.8736
epsilon[, 2] <- (rchisq(n, 5) + epsilon_z_y) * (sqrt(3/20)) - 3.8736
# Simulate selection equation
z_star <- 1 + 1 * X[,1] + 1 * X[,2] + epsilon[,1]
z <- as.numeric((z_star > 0))

# Simulate outcome equation
y_star <- 1 + 1 * X[,1] + 1 * X[,3] + epsilon[,2]
z <- as.numeric((z_star > 0))
y <- y_star
y[z==0] <- NA
h <- as.data.frame(cbind(z, y, X))
names(h) <- c("z", "y", "x1", "x2", "x3")

# Estimate parameters
model <- hpaSelection(selection = z ~ x1 + x2,
                     outcome = y ~ x1 + x3,
                     data = h, z_K = 1, y_K = 3)

summary(model)

# Get conditional predictions for outcome equation
model_pred_c <- predict(model, is_cond = TRUE)
# Conditional predictions y|z=1
model_pred_c$y_1
# Conditional predictions y|z=0
model_pred_c$y_0

# Get unconditional predictions for outcome equation
model_pred_u <- predict(model, is_cond = FALSE)
model_pred_u$y

```

```
# Get conditional predictions for selection equation
# Note that for z=0 these predictions are NA
predict(model, is_cond = TRUE, is_outcome = FALSE)
# Get unconditional predictions for selection equation
predict(model, is_cond = FALSE, is_outcome = FALSE)
```

ihpa

Interval distribution function hermite polynomial approximation

Description

This function calculates interval distribution function hermite polynomial approximation.

Usage

```
ihpa(
  x_lower = matrix(1, 1),
  x_upper = matrix(1, 1),
  pol_coefficients = numeric(0),
  pol_degrees = numeric(0),
  given_ind = logical(0),
  omit_ind = logical(0),
  mean = numeric(0),
  sd = numeric(0),
  is_parallel = FALSE,
  is_log = FALSE
)
```

Arguments

<code>x_lower</code>	numeric matrix of lower integration limits. Note that <code>x_lower</code> rows are observations while variables are columns.
<code>x_upper</code>	numeric matrix of upper integration limits. Note that <code>x_upper</code> rows are observations while variables are columns.
<code>pol_coefficients</code>	numeric vector of polynomial coefficients.
<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
<code>given_ind</code>	logical vector indicating wheather corresponding component is conditioned. By default it is a logical vector of FALSE values.
<code>omit_ind</code>	logical vector indicating wheather corresponding component is omitted. By default it is a logical vector of FALSE values.
<code>mean</code>	numeric vector of expected values.

<code>sd</code>	positive numeric vector of standard deviations.
<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
<code>is_log</code>	logical; if TRUE then probabilities <code>p</code> are given as $\log(p)$ or derivatives will be given respect to $\log(p)$

Details

Interval distribution function represents probability that random vector components will be greater then values given in `x_lower` and lower then values that are in `x_upper`.

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters `mean` and `sd` determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below.

Parameters `mean`, `sd`, `given_ind`, `omit_ind` should have the same length as `pol_degrees` parameter.

Value

This function returns interval distribution function hermite polynomial approximation at point `x`.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

Examples

```
## Let's approximate some three random variables joint interval distribution
## function (idf) at lower and upper points (0.1, 0.2, 0.3) and
## (0.4, 0.5, 0.6) correspondingly with hermite polynomial of (1,2,3) degrees
## which polynomial coefficients equals 1 except coefficient related to
## x1*(x^3) polynomial element which equals 2. Also suppose that normal
## density related mean vector equals (1.1, 1.2, 1.3) while standard
## deviations vector is (2.1, 2.2, 2.3).

## Prepare initial values
x_lower <- matrix(c(0.1, 0.2, 0.3), nrow=1)
x_upper <- matrix(c(0.4, 0.5, 0.6), nrow=1)
mean <- c(1.1, 1.2, 1.3)
sd <- c(2.1, 2.2, 2.3)
pol_degrees <- c(1, 2, 3)

# Create polynomial powers and indexes correspondence matrix
pol_ind <- polynomialIndex(pol_degrees)

# Set all polynomial coefficients to 1
pol_coefficients <- rep(1, ncol(pol_ind))
```

```

pol_degrees_n <- length(pol_degrees)

# Assign coefficient 2 to the polynomial element(x1 ^ 1)*(x2 ^ 0)*(x3 ^ 2)
pol_coefficients[apply(pol_ind, 2, function(x) all(x == c(1, 0, 2)))] <- 2

# Visualize correspondence between polynomial
# elements and their coefficients
as.data.frame(rbind(pol_ind, pol_coefficients),
row.names = c("x1 power", "x2 power", "x3 power", "coefficients"),
optional = TRUE)
printPolynomial(pol_degrees, pol_coefficients)

# Calculate idf approximation at points x_lower and x_upper
ihpa(x_lower = x_lower, x_upper = x_upper,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd)

# Condition second component to be 0.7
# Substitute x second component with conditional value 0.7
x_upper <- matrix(c(0.4, 0.7, 0.6), nrow = 1)

# Set TRUE to the second component indicating that it is conditioned
given_ind <- c(FALSE, TRUE, FALSE)

# Calculate conditional(on x2 = 0.5) idf approximation at points x_lower and x_upper
ihpa(x_lower = x_lower, x_upper = x_upper,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
given_ind = given_ind)

# Consider third component marginal distribution
# conditioned on the second component 0.7 value
# Set TRUE to the first component indicating that it is omitted
omit_ind <- c(TRUE, FALSE, FALSE)

# Calculate conditional (on x2=0.5) marginal (for x3)
# idf approximation at points x_lower and x_upper
ihpa(x_lower = x_lower, x_upper = x_upper,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
given_ind = given_ind, omit_ind = omit_ind)

```

ihpaDiff

Calculate gradient of interval distribution function hermite polynomial approximation

Description

This function calculates gradient of interval distribution function hermite polynomial approximation.

Usage

```
ihpaDiff(
  x_lower = matrix(1, 1),
  x_upper = matrix(1, 1),
  pol_coefficients = numeric(0),
  pol_degrees = numeric(0),
  given_ind = logical(0),
  omit_ind = logical(0),
  mean = numeric(0),
  sd = numeric(0),
  type = "pol_coefficients",
  is_parallel = FALSE,
  is_log = FALSE
)
```

Arguments

<code>x_lower</code>	numeric matrix of lower integration limits. Note that <code>x_lower</code> rows are observations while variables are columns.
<code>x_upper</code>	numeric matrix of upper integration limits. Note that <code>x_upper</code> rows are observations while variables are columns.
<code>pol_coefficients</code>	numeric vector of polynomial coefficients.
<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
<code>given_ind</code>	logical vector indicating wheather corresponding component is conditioned. By default it is a logical vector of FALSE values.
<code>omit_ind</code>	logical vector indicating wheather corresponding component is omitted. By default it is a logical vector of FALSE values.
<code>mean</code>	numeric vector of expected values.
<code>sd</code>	positive numeric vector of standard deviations.
<code>type</code>	determines the partial derivatives to be included into gradient. If <code>type="pol_coefficients"</code> then gradient will contain partial derivatives respect to polynomial coefficients listed in the same order as <code>pol_coefficients</code> . Other available types are <code>type="mean"</code> and <code>type="sd"</code> . For function dhpaDiff it is possible to take gradient respect to the x points setting <code>type="x"</code> . For function ihpaDiff it is possible to take gradient respect to the x lower and upper points setting <code>type="x_lower"</code> or <code>type="upper"</code> correspondingly. In order to get full gradient please set <code>type="all"</code> .
<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
<code>is_log</code>	logical; if TRUE then probabilities p are given as log(p) or derivatives will be given respect to log(p)

Details

Interval distribution function represents probability that random vector components will be greater then values given in `x_lower` and lower then values that are in `x_upper`.

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters `mean` and `sd` determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below.

Parameters `mean`, `sd`, `given_ind`, `omit_ind` should have the same length as `pol_degrees` parameter.

If `x` has more then one row then the output will be jacobian matrix where rows are gradients.

Value

This function returns gradient of interval distribution function hermite polynomial approximation at point `x`. Gradient elements are determined by the `type` argument.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

Examples

```
## Let's approximate some three random variables joint interval distribution
## function (idf) at lower and upper points (0.1, 0.2, 0.3) and
## (0.4, 0.5, 0.6) correspondingly with hermite polynomial of (1,2,3)
## degrees which polynomial coefficients equals 1 except coefficient
## related to x1*(x^3) polynomial element which equals 2.
## Also suppose that normal density related mean vector equals
## (1.1, 1.2, 1.3) while standard deviations vector is (2.1, 2.2, 2.3).
## In this example let's calculate interval distribution approximating
## function gradient respect to polynomial coefficients.

# Prepare initial values
x_lower <- matrix(c(0.1, 0.2, 0.3), nrow=1)
x_upper <- matrix(c(0.4, 0.5, 0.6), nrow=1)
mean <- c(1.1, 1.2, 1.3)
sd <- c(2.1, 2.2, 2.3)
pol_degrees <- c(1, 2, 3)

# Create polynomial powers and indexes correspondence matrix
pol_ind <- polynomialIndex(pol_degrees)

# Set all polynomial coefficients to 1
pol_coefficients <- rep(1, ncol(pol_ind))
pol_degrees_n <- length(pol_degrees)

# Assign coefficient 2 to the polynomial element(x1 ^ 1)*(x2 ^ 0)*(x3 ^ 2)
pol_coefficients[apply(pol_ind, 2, function(x) all(x == c(1, 0, 2)))] <- 2
```

```

# Visualize correspondence between polynomial
# elements and their coefficients
as.data.frame(rbind(pol_ind, pol_coefficients),
  row.names = c("x1 power", "x2 power", "x3 power", "coefficients"),
  optional = TRUE)
printPolynomial(pol_degrees, pol_coefficients)

# Calculate idf approximation gradient respect to
# polynomial coefficients at points x_lower and x_upper
ihpaDiff(x_lower = x_lower, x_upper = x_upper,
  pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
  mean = mean, sd = sd)

# Condition second component to be 0.7
# Substitute x second component with conditional value 0.7
x_upper <- matrix(c(0.4, 0.7, 0.6), nrow = 1)

# Set TRUE to the second component indicating that it is conditioned
given_ind <- c(FALSE, TRUE, FALSE)

# Calculate conditional(on x2 = 0.5) idf approximation
# respect to polynomial coefficients at points x_lower and x_upper
ihpaDiff(x_lower = x_lower, x_upper = x_upper,
  pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
  mean = mean, sd = sd,
  given_ind = given_ind)

# Consider third component marginal distribution
# conditioned on the second component 0.7 value
# Set TRUE to the first component indicating that it is omitted
omit_ind <- c(TRUE, FALSE, FALSE)

# Calculate conditional (on x2=0.5) marginal (for x3) idf approximation
# respect to:
# polynomial coefficients
ihpaDiff(x_lower = x_lower, x_upper = x_upper,
  pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
  mean = mean, sd = sd,
  given_ind = given_ind, omit_ind = omit_ind)
# mean
ihpaDiff(x_lower = x_lower, x_upper = x_upper,
  pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
  mean = mean, sd = sd,
  given_ind = given_ind, omit_ind = omit_ind,
  type = "mean")
# sd
ihpaDiff(x_lower = x_lower, x_upper = x_upper,
  pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
  mean = mean, sd = sd,
  given_ind = given_ind, omit_ind = omit_ind,
  type = "sd")
# x_lower

```

```

ihpaDiff(x_lower = x_lower, x_upper = x_upper,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
given_ind = given_ind, omit_ind = omit_ind,
type = "x_lower")
# x_upper
ihpaDiff(x_lower = x_lower, x_upper = x_upper,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
given_ind = given_ind, omit_ind = omit_ind,
type = "x_upper")

```

itrhpa	<i>Truncated interval distribution function hermite polynomial approximation for truncated distribution</i>
--------	---

Description

This function calculates truncated interval distribution function hermite polynomial approximation for truncated distribution.

Usage

```

itrhpa(
  x_lower = matrix(1, 1),
  x_upper = matrix(1, 1),
  tr_left = matrix(1, 1),
  tr_right = matrix(1, 1),
  pol_coefficients = numeric(0),
  pol_degrees = numeric(0),
  given_ind = logical(0),
  omit_ind = logical(0),
  mean = numeric(0),
  sd = numeric(0),
  is_parallel = FALSE,
  is_log = FALSE
)

```

Arguments

<code>x_lower</code>	numeric matrix of lower integration limits. Note that <code>x_lower</code> rows are observations while variables are columns.
<code>x_upper</code>	numeric matrix of upper integration limits. Note that <code>x_upper</code> rows are observations while variables are columns.
<code>tr_left</code>	numeric matrix of left (lower) truncation limits. Note that <code>tr_right</code> rows are observations while variables are columns. If <code>tr_left</code> or <code>tr_right</code> is single row matrix then the same truncation limits would be applied to all observations that are determined by the first rows of these matrices.

<code>tr_right</code>	numeric matrix of right (upper) truncation limits. Note that <code>tr_right</code> rows are observations while variables are columns. If <code>tr_left</code> or <code>tr_right</code> is single row matrix then the same truncation limits would be applied to all observations that are determined by the first rows of these matrices.
<code>pol_coefficients</code>	numeric vector of polynomial coefficients.
<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
<code>given_ind</code>	logical vector indicating wheather corresponding component is conditioned. By default it is a logical vector of FALSE values.
<code>omit_ind</code>	logical vector indicating wheather corresponding component is omitted. By default it is a logical vector of FALSE values.
<code>mean</code>	numeric vector of expected values.
<code>sd</code>	positive numeric vector of standard deviations.
<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
<code>is_log</code>	logical; if TRUE then probabilities p are given as $\log(p)$ or derivatives will be given respect to $\log(p)$

Details

Interval distribution function represents probability that random vector components will be greater then values given in `x_lower` and lower then values that are in `x_upper`.

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters `mean` and `sd` determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below.

Parameters `mean`, `sd`, `given_ind`, `omit_ind` should have the same length as `pol_degrees` parameter.

Value

This function returns interval distribution function (`idf`) hermite polynomial approximation at point `x` for truncated distribution.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

Examples

```
## Let's approximate some three truncated random variables joint interval
## distribution function at lower and upper points (0,1, 0.2, 0.3) and
## (0.4, 0.5, 0.6) correspondingly with hermite polynomial of (1,2,3)
## degrees which polynomial coefficients equals 1 except coefficient
```

```

## related to  $x_1 \cdot (x^3)$  polynomial element which equals 2. Also suppose
## that normal density related mean vector equals (1.1, 1.2, 1.3) while
## standard deviations vector is (2.1, 2.2, 2.3). Suppose that lower and
## upper truncation are (-1.1,-1.2,-1.3) and (1.1,1.2,1.3) correspondingly.

# Prepare initial values
x_lower <- matrix(c(0.1, 0.2, 0.3), nrow=1)
x_upper <- matrix(c(0.4, 0.5, 0.6), nrow=1)
tr_left = matrix(c(-1.1,-1.2,-1.3), nrow = 1)
tr_right = matrix(c(1.1,1.2,1.3), nrow = 1)
mean <- c(1.1, 1.2, 1.3)
sd <- c(2.1, 2.2, 2.3)
pol_degrees <- c(1, 2, 3)

# Create polynomial powers and indexes correspondence matrix
pol_ind <- polynomialIndex(pol_degrees)
# Set all polynomial coefficients to 1
pol_coefficients <- rep(1, ncol(pol_ind))
pol_degrees_n <- length(pol_degrees)

# Assign coefficient 2 to the polynomial element  $(x_1^1) \cdot (x_2^0) \cdot (x_3^2)$ 
pol_coefficients[apply(pol_ind, 2, function(x) all(x == c(1, 0, 2)))] <- 2
# Visualize correspondence between polynomial
# elements and their coefficients
as.data.frame(rbind(pol_ind, pol_coefficients),
row.names = c("x1 power", "x2 power", "x3 power", "coefficients"),
optional = TRUE)
printPolynomial(pol_degrees, pol_coefficients)

# Calculate idf approximation at points x_lower and x_upper
itrhpa(x_lower = x_lower, x_upper = x_upper,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
tr_left = tr_left, tr_right = tr_right)

# Condition second component to be 0.7
# Substitute x second component with conditional value 0.7
x_upper <- matrix(c(0.4, 0.7, 0.6), nrow = 1)
# Set TRUE to the second component indicating that it is conditioned
given_ind <- c(FALSE, TRUE, FALSE)

# Calculate conditional(on  $x_2 = 0.5$ ) idf
# approximation at points x_lower and x_upper
itrhpa(x_lower = x_lower, x_upper = x_upper,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
given_ind = given_ind,
tr_left = tr_left, tr_right = tr_right)

# Consider third component marginal distribution
# conditioned on the second component 0.7 value
# Set TRUE to the first component indicating that it is omitted
omit_ind <- c(TRUE, FALSE, FALSE)

```

```
# Calculate conditional (on x2=0.5) marginal (for x3) idf
# approximation at points x_lower and x_upper
itrhpa(x_lower = x_lower, x_upper = x_upper,
       pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
       mean = mean, sd = sd,
       given_ind = given_ind, omit_ind = omit_ind,
       tr_left = tr_left, tr_right = tr_right)
```

logLik.hpaBinary	<i>Calculates log-likelihood for "hpaBinary" object</i>
------------------	---

Description

This function calculates log-likelihood for "hpaBinary" object

Usage

```
## S3 method for class 'hpaBinary'
logLik(object, ...)
```

Arguments

object	Object of class "hpaBinary"
...	further arguments (currently ignored)

logLik.hpaML	<i>Calculates log-likelihood for "hpaML" object</i>
--------------	---

Description

This function calculates log-likelihood for "hpaML" object

Usage

```
## S3 method for class 'hpaML'
logLik(object, ...)
```

Arguments

object	Object of class "hpaML"
...	further arguments (currently ignored)

logLik.hpaSelection	<i>Calculates log-likelihood for "hpaSelection" object</i>
---------------------	--

Description

This function calculates log-likelihood for "hpaSelection" object

Usage

```
## S3 method for class 'hpaSelection'  
logLik(object, ...)
```

Arguments

object	Object of class "hpaSelection"
...	further arguments (currently ignored)

logLik_hpaBinary	<i>Calculates log-likelihood for "hpaBinary" object</i>
------------------	---

Description

This function calculates log-likelihood for "hpaBinary" object

Usage

```
logLik_hpaBinary(object)
```

Arguments

object	Object of class "hpaBinary"
--------	-----------------------------

logLik_hpaML	<i>Calculates log-likelihood for "hpaML" object</i>
--------------	---

Description

This function calculates log-likelihood for "hpaML" object

Usage

```
logLik_hpaML(object)
```

Arguments

object	Object of class "hpaML"
--------	-------------------------

logLik_hpaSelection	<i>Calculates log-likelihood for "hpaSelection" object</i>
---------------------	--

Description

This function calculates log-likelihood for "hpaSelection" object

Usage

```
logLik_hpaSelection(object)
```

Arguments

object	Object of class "hpaSelection"
--------	--------------------------------

mecdf	<i>Calculates multivariate empirical cumulative distribution function</i>
-------	---

Description

This function calculates multivariate empirical cumulative distribution function at each point of the sample

Usage

```
mecdf(x)
```

Arguments

x	numeric matrix which rows are observations
---	--

normalMoment	<i>Calculate k-th order moment of normal distribution</i>
--------------	---

Description

This function recursively calculates k-th order moment of normal distribution.

Usage

```
normalMoment(
  k = 0L,
  mean = 0,
  sd = 1,
  return_all_moments = FALSE,
  is_validation = TRUE,
  is_central = FALSE,
  diff_type = "NO"
)
```

Arguments

<code>k</code>	non-negative integer moment order.
<code>mean</code>	numeric expected value.
<code>sd</code>	positive numeric standard deviation.
<code>return_all_moments</code>	logical; if TRUE, function returns (k+1)-dimensional numeric vector of moments of normally distributed random variable with mean = mean and standard deviation = sd. Note that i-th vector's component value corresponds to the (i-1)-th moment.
<code>is_validation</code>	logical value indicating whether function input arguments should be validated. Set it to FALSE for slight performance boost (default value is TRUE).
<code>is_central</code>	logical; if TRUE, then central moments will be calculated.
<code>diff_type</code>	string indicating the type of the argument the moment should be differentiated respect to before return. Default value is "NO" so the moments itself will be returned. Alternative values are "mean", "sd", x_lower and x_upper.

Details

This function estimates k-th order moment of normal distribution which mean equals to mean and standard deviation equals to sd.

Note that parameter k value automatically converts to integer. So passing non-integer k value will not cause any errors but the calculations will be performed for rounded k value only.

Value

This function returns k-th order moment of normal distribution which mean equals to mean and standard deviation is sd. If return_all_moments is TRUE then see this argument description above for output details.

Examples

```
## Calculate 5-th order moment of normal random variable which
## mean equals to 3 and standard deviation is 5.
```

```
# 5-th moment
normalMoment(k = 5, mean = 3, sd = 5)

# (0-5)-th moments
normalMoment(k = 5, mean = 3, sd = 5, return_all_moments = TRUE)

# 5-th moment derivative respect to mean
normalMoment(k = 5, mean = 3, sd = 5, diff_type = "mean")

# 5-th moment derivative respect to sd
normalMoment(k = 5, mean = 3, sd = 5, diff_type = "sd")
```

phpa

Distribution function hermite polynomial approximation

Description

This function calculates cumulative distribution function hermite polynomial approximation.

Usage

```
phpa(
  x = matrix(1, 1),
  pol_coefficients = numeric(0),
  pol_degrees = numeric(0),
  given_ind = logical(0),
  omit_ind = logical(0),
  mean = numeric(0),
  sd = numeric(0),
  is_parallel = FALSE,
  is_log = FALSE
)
```

Arguments

<code>x</code>	numeric matrix of cumulative distribution function arguments. Note that <code>x</code> rows are observations while variables are columns.
<code>pol_coefficients</code>	numeric vector of polynomial coefficients.
<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
<code>given_ind</code>	logical vector indicating wheather corresponding component is conditioned. By default it is a logical vector of FALSE values.
<code>omit_ind</code>	logical vector indicating wheather corresponding component is omitted. By default it is a logical vector of FALSE values.
<code>mean</code>	numeric vector of expected values.
<code>sd</code>	positive numeric vector of standard deviations.

<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
<code>is_log</code>	logical; if TRUE then probabilities <code>p</code> are given as <code>log(p)</code> or derivatives will be given respect to <code>log(p)</code>

Details

Densities hermite polynomial approximation approach has been proposed by A. Gallant and D. W. Nychka in 1987. The main idea is to approximate unknown distribution density with hermite polynomial of degree `pol_degree`. In this framework hermite polynomial represents adjusted (to insure integration to 1) product of squared polynomial and normal distribution densities. Parameters `mean` and `sd` determine means and standard deviations of normal distribution density functions which are parts of this polynomial. For more information please refer to the literature listed below. Parameters `mean`, `sd`, `given_ind`, `omit_ind` should have the same length as `pol_degrees` parameter.

Value

This function returns cumulative distribution function hermite polynomial approximation at point `x`.

References

A. Gallant and D. W. Nychka (1987) <doi:10.2307/1913241>

Examples

```
## Let's approximate some three random variables joint cumulative
## distribution function (cdf) at point (0.1, 0.2, 0.3)
## with hermite polynomial of (1,2,3) degrees which polynomial
## coefficients equals 1 except coefficient related to x1*(x^3) polynomial
## element which equals 2. Also suppose that normal density related
## mean vector equals (1.1, 1.2, 1.3) while standard deviations
## vector is (2.1, 2.2, 2.3).

## Prepare initial values
x <- matrix(c(0.1, 0.2, 0.3), nrow=1)
mean <- c(1.1, 1.2, 1.3)
sd <- c(2.1, 2.2, 2.3)
pol_degrees <- c(1, 2, 3)

# Create polynomial powers and indexes correspondence matrix
pol_ind <- polynomialIndex(pol_degrees)

# Set all polynomial coefficients to 1
pol_coefficients <- rep(1, ncol(pol_ind))
pol_degrees_n <- length(pol_degrees)

# Assign coefficient 2 to the polynomial element(x1 ^ 1)*(x2 ^ 0)*(x3 ^ 2)
pol_coefficients[apply(pol_ind, 2, function(x) all(x == c(1, 0, 2)))] <- 2
```

```

# Visualize correspondence between polynomial
# elements and their coefficients
as.data.frame(rbind(pol_ind, pol_coefficients),
row.names = c("x1 power", "x2 power", "x3 power", "coefficients"),
optional = TRUE)
printPolynomial(pol_degrees, pol_coefficients)

# Calculate cdf approximation at point x
phpa(x = x,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd)

# Condition second component to be 0.5
# Substitute x second component with conditional value 0.5
x <- matrix(c(0.1, 0.5, 0.3), nrow = 1)

# Set TRUE to the second component indicating that it is conditioned
given_ind <- c(FALSE, TRUE, FALSE)

# Calculate conditional(on x2 = 0.5) cdf approximation at point x
phpa(x = x,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
given_ind = given_ind)

# Consider third component marginal distribution
# conditioned on the second component 0.5 value

# Set TRUE to the first component indicating that it is omitted
omit_ind <- c(TRUE, FALSE, FALSE)

# Calculate conditional (on x2=0.5) marginal (for x3) cdf approximation at point x
phpa(x = x,
pol_coefficients = pol_coefficients, pol_degrees = pol_degrees,
mean = mean, sd = sd,
given_ind = given_ind, omit_ind = omit_ind)

```

plot.hpaBinary

Plot hpaBinary random errors approximated density

Description

Plot hpaBinary random errors approximated density

Usage

```

## S3 method for class 'hpaBinary'
plot(x, y = NULL, ...)

```

Arguments

x	Object of class "hpaBinary"
y	this parameter currently ignored
...	further arguments (currently ignored)

plot.hpaML

*Plot approximated marginal density using hpaML output***Description**

Plot approximated marginal density using hpaML output

Usage

```
## S3 method for class 'hpaML'
plot(x, y = NULL, ..., ind = 1, given = NULL)
```

Arguments

x	Object of class "hpaML"
y	this parameter currently ignored
...	further arguments (currently ignored)
ind	= index of random variable for which approximation to marginal density should be plotted
given	= numeric vector of the same length as given_ind from x. Determines conditional values for the corresponding components. NA values in given vector indicate that corresponding random variable is not conditioned. By default all given components are NA so unconditional marginal density will be plotted for the ind-the random variable.

plot.hpaSelection

*Plot hpaSelection random errors approximated density***Description**

Plot hpaSelection random errors approximated density

Usage

```
## S3 method for class 'hpaSelection'
plot(x, y = NULL, ..., is_outcome = TRUE)
```

Arguments

x	Object of class "hpaSelection"
y	this parameter currently ignored
...	further arguments (currently ignored)
is_outcome	logical; if TRUE then function plots the graph for outcome equation random errors. Otherwise plot for selection equation random errors will be plotted.

Value

This function returns the list containing random error's expected value errors_exp and variance errors_var estimates for selection (if is_outcome = TRUE) or outcome (if is_outcome = FALSE) equation.

plot_hpaBinary	<i>Plot hpaBinary random errors approximated density</i>
----------------	--

Description

Plot hpaBinary random errors approximated density

Usage

```
plot_hpaBinary(x)
```

Arguments

x	Object of class "hpaBinary"
---	-----------------------------

plot_hpaSelection	<i>Plot hpaSelection random errors approximated density</i>
-------------------	---

Description

Plot hpaSelection random errors approximated density

Usage

```
plot_hpaSelection(x, is_outcome = TRUE)
```

Arguments

x	Object of class "hpaSelection"
is_outcome	logical; if TRUE then function plots the graph for outcome equation random errors. Otherwise plot for selection equation random errors will be plotted.

Value

This function returns the list containing random error's expected value `errors_exp` and variance `errors_var` estimates for selection (if `is_outcome = TRUE`) or outcome (if `is_outcome = FALSE`) equation.

<code>pnorm_parallel</code>	<i>Calculate normal cdf in parallel</i>
-----------------------------	---

Description

Calculate in parallel for each value from vector `x` distribution function of normal distribution with mean equal to mean and standard deviation equal to `sd`.

Usage

```
pnorm_parallel(x, mean = 0, sd = 1, is_parallel = FALSE)
```

Arguments

<code>x</code>	vector of quantiles: should be numeric vector, not just double value.
<code>mean</code>	double value.
<code>sd</code>	double positive value.
<code>is_parallel</code>	if <code>TRUE</code> then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).

<code>polynomialIndex</code>	<i>Returns matrix of polynomial indexes</i>
------------------------------	---

Description

Returns matrix of polynomial indexes for the polynomial with degrees (orders) vector `pol_degrees`.

Usage

```
polynomialIndex(pol_degrees = 0L)
```

Arguments

<code>pol_degrees</code>	non-negative integer vector of polynomial degrees.
--------------------------	--

Details

This function motivation is to have an opportunity to iterate through the columns of polynomial indexes matrix in order to access polynomial elements being aware of their powers.

Value

This function returns polynomial indexes matrix which rows are responsible for variables while columns are related to powers.

Examples

```
## Get polynomial indexes matrix for the polynomial
## which degrees are (1, 3, 5)

polynomialIndex(c(1, 3, 5))

## Consider polynomial of degrees (2, 1) such that coefficients
## for elements which powers sum is even are 2 and for those which powers
## are odd are 5. So the polynomial is  $2+5y+5x+2xy+2x^2+5yx^2$ .

# Let's represent its powers (not coefficients) in a matrix form
pol_matrix <- polynomialIndex(c(2, 1))

# Suppose we want to calculate this polynomial coefficients sum:
powers_sum <- 0

# For pedagogical reasons iterate through the pol_matrix columns
pol_matrix_length = dim(pol_matrix)[2]

for (i in 1:pol_matrix_length)
{
  if ((pol_matrix[1, i] + pol_matrix[2, i]) %% 2 == 0)
  {
    powers_sum <- powers_sum + 2
  } else {
    powers_sum <- powers_sum + 5
  }
}
# powers_sum value will be 21
```

predict.hpaBinary	<i>Predict method for hpaBinary</i>
-------------------	-------------------------------------

Description

Predict method for hpaBinary

Usage

```
## S3 method for class 'hpaBinary'
predict(object, ..., newdata = NULL, is_prob = TRUE)
```

Arguments

object	Object of class "hpaBinary"
...	further arguments (currently ignored)
newdata	An optional data frame (for hpaBinary and hpaSelection) or numeric matrix (for hpaML) in which to look for variables with which to predict. If omitted, the original dataframe (matrix) used.
is_prob	logical; if TRUE (default) then function returns predicted probabilities. Otherwise latent variable (single index) estimates will be returned.

Value

This function returns predicted probabilities based on [hpaBinary](#) estimation results.

predict.hpaML	<i>Predict method for hpaML</i>
---------------	---------------------------------

Description

Predict method for hpaML

Usage

```
## S3 method for class 'hpaML'
predict(object, ..., newdata = matrix(c(0)))
```

Arguments

object	Object of class "hpaML"
...	further arguments (currently ignored)
newdata	An optional data frame (for hpaBinary and hpaSelection) or numeric matrix (for hpaML) in which to look for variables with which to predict. If omitted, the original dataframe (matrix) used.

Value

This function returns predictions based on [hpaML](#) estimation results.

predict.hpaSelection *Predict outcome and selection equation values from hpaSelection model*

Description

This function predicts outcome and selection equation values from hpaSelection model.

Usage

```
## S3 method for class 'hpaSelection'
predict(
  object,
  ...,
  newdata = NULL,
  method = "HPA",
  is_cond = TRUE,
  is_outcome = TRUE
)
```

Arguments

object	Object of class "hpaSelection"
...	further arguments (currently ignored)
newdata	An optional data frame (for hpaBinary and hpaSelection) or numeric matrix (for hpaML) in which to look for variables with which to predict. If omitted, the original dataframe (matrix) used.
method	string value indicating prediction method based on hermite polynomial approximation "HPA" or Newey method "Newey".
is_cond	logical; if TRUE (default) then conditional predictions will be estimated. Otherwise unconditional predictions will be returned.
is_outcome	logical; if TRUE (default) then predictions for selection equation will be estimated using "HPA" method. Otherwise selection equation predictions (probabilities) will be returned.

Details

Note that Newey method can't predict conditional outcomes for zero selection equation value. Conditional probabilities for selection equation could be estimated only when dependent variable from outcome equation is observable.

Value

This function returns the list which structure depends on method, is_probit and is_outcome values.

predict_hpaBinary	<i>Predict method for hpaBinary</i>
-------------------	-------------------------------------

Description

Predict method for hpaBinary

Usage

```
predict_hpaBinary(object, newdata = NULL, is_prob = TRUE)
```

Arguments

object	Object of class "hpaBinary"
newdata	An optional data frame (for hpaBinary and hpaSelection) or numeric matrix (for hpaML) in which to look for variables with which to predict. If omitted, the original dataframe (matrix) used.
is_prob	logical; if TRUE (default) then function returns predicted probabilities. Otherwise latent variable (single index) estimates will be returned.

Value

This function returns predicted probabilities based on [hpaBinary](#) estimation results.

predict_hpaML	<i>Predict method for hpaML</i>
---------------	---------------------------------

Description

Predict method for hpaML

Usage

```
predict_hpaML(object, newdata = matrix(1, 1))
```

Arguments

object	Object of class "hpaML"
newdata	An optional data frame (for hpaBinary and hpaSelection) or numeric matrix (for hpaML) in which to look for variables with which to predict. If omitted, the original dataframe (matrix) used.

Value

This function returns predictions based on [hpaML](#) estimation results.

predict_hpaSelection	<i>Predict outcome and selection equation values from hpaSelection model</i>
----------------------	--

Description

This function predicts outcome and selection equation values from hpaSelection model.

Usage

```
predict_hpaSelection(
  object,
  newdata = NULL,
  method = "HPA",
  is_cond = TRUE,
  is_outcome = TRUE
)
```

Arguments

object	Object of class "hpaSelection"
newdata	An optional data frame (for hpaBinary and hpaSelection) or numeric matrix (for hpaML) in which to look for variables with which to predict. If omitted, the original dataframe (matrix) used.
method	string value indicating prediction method based on hermite polynomial approximation "HPA" or Newey method "Newey".
is_cond	logical; if TRUE (default) then conditional predictions will be estimated. Otherwise unconditional predictions will be returned.
is_outcome	logical; if TRUE (default) then predictions for selection equation will be estimated using "HPA" method. Otherwise selection equation predictions (probabilities) will be returned.

Details

Note that Newey method can't predict conditional outcomes for zero selection equation value. Conditional probabilities for selection equation could be estimated only when dependent variable from outcome equation is observable.

Value

This function returns the list which structure depends on method, is_probit and is_outcome values.

print.hpaBinary	<i>Print method for "hpaBinary" object</i>
-----------------	--

Description

Print method for "hpaBinary" object

Usage

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

Arguments

x	Object of class "hpaBinary"
...	further arguments (currently ignored)

print.hpaML	<i>Print method for "hpaML" object</i>
-------------	--

Description

Print method for "hpaML" object

Usage

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

Arguments

x	Object of class "hpaML"
...	further arguments (currently ignored)

print.hpaSelection	<i>Print method for "hpaSelection" object</i>
--------------------	---

Description

Print method for "hpaSelection" object

Usage

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

Arguments

x	Object of class "hpaSelection"
...	further arguments (currently ignored)

print.summary.hpaBinary	<i>Summary for "hpaBinary" object</i>
-------------------------	---------------------------------------

Description

Summary for "hpaBinary" object

Usage

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

Arguments

x	Object of class "hpaBinary"
...	further arguments (currently ignored)

print.summary.hpaML *Summary for hpaML output*

Description

Summary for hpaML output

Usage

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

Arguments

x	Object of class "hpaML"
...	further arguments (currently ignored)

print.summary.hpaSelection
 Summary for "hpaSelection" object

Description

Summary for "hpaSelection" object

Usage

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

Arguments

x	Object of class "hpaSelection"
...	further arguments (currently ignored)

printPolynomial	<i>Print polynomial given it's degrees and coefficients</i>
-----------------	---

Description

This function prints polynomial given it's degrees and coefficients.

Usage

```
printPolynomial(pol_degrees, pol_coefficients)
```

Arguments

`pol_degrees` non-negative integer vector of polynomial degrees.
`pol_coefficients` numeric vector of polynomial coefficients.

Details

Function automatically removes polynomial elements which coefficient are zero and variables which power is zero. Output may contain long coefficients representation as they are not rounded.

Value

This function returns the string which contains polynomial symbolic representation.

Examples

```
## Let's represent polynomial 0.3+0.5x2-x2^2+2x1+1.5x1x2+x1x2^2

pol_degrees <- c(1, 2)
pol_coefficients <- c(0.3, 0.5, -1, 2, 1.5, 1)

printPolynomial(pol_degrees, pol_coefficients)
```

print_summary_hpaBinary	<i>Summary for hpaBinary output</i>
-------------------------	-------------------------------------

Description

Summary for hpaBinary output

Usage

```
print_summary_hpaBinary(x)
```

Arguments

x Object of class "hpaML"

print_summary_hpaML *Summary for hpaML output*

Description

Summary for hpaML output

Usage

print_summary_hpaML(x)

Arguments

x Object of class "hpaML"

print_summary_hpaSelection
 Summary for hpaSelection output

Description

Summary for hpaSelection output

Usage

print_summary_hpaSelection(x)

Arguments

x Object of class "hpaSelection"

summary.hpaBinary	<i>Summarizing hpaBinary Fits</i>
-------------------	-----------------------------------

Description

Summarizing hpaBinary Fits

Usage

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

Arguments

object	Object of class "hpaBinary"
...	further arguments (currently ignored)

Value

This function returns the same list as [hpaBinary](#) function changing it's class to "summary.hpaBinary".

summary.hpaML	<i>Summarizing hpaML Fits</i>
---------------	-------------------------------

Description

Summarizing hpaML Fits

Usage

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

Arguments

object	Object of class "hpaML"
...	further arguments (currently ignored)

Value

This function returns the same list as [hpaML](#) function changing it's class to "summary.hpaML".

summary.hpaSelection	<i>Summarizing hpaSelection Fits</i>
----------------------	--------------------------------------

Description

This function summarizing hpaSelection Fits

Usage

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

Arguments

object	Object of class "hpaSelection"
...	further arguments (currently ignored)

Value

This function returns the same list as [hpaSelection](#) function changing it's class to "summary.hpaSelection".

summary_hpaBinary	<i>Summarizing hpaBinary Fits</i>
-------------------	-----------------------------------

Description

Summarizing hpaBinary Fits

Usage

```
summary_hpaBinary(object)
```

Arguments

object	Object of class "hpaBinary"
--------	-----------------------------

Value

This function returns the same list as [hpaBinary](#) function changing it's class to "summary.hpaBinary".

summary_hpaML	<i>Summarizing hpaML Fits</i>
---------------	-------------------------------

Description

Summarizing hpaML Fits

Usage

```
summary_hpaML(object)
```

Arguments

object	Object of class "hpaML"
--------	-------------------------

Value

This function returns the same list as [hpaML](#) function changing it's class to "summary.hpaML".

summary_hpaSelection	<i>Summarizing hpaSelection Fits</i>
----------------------	--------------------------------------

Description

This function summarizing hpaSelection Fits

Usage

```
summary_hpaSelection(object)
```

Arguments

object	Object of class "hpaSelection"
--------	--------------------------------

Value

This function returns the same list as [hpaSelection](#) function changing it's class to "summary.hpaSelection".

truncatedNormalMoment *Calculate k-th order moment of truncated normal distribution*

Description

This function recursively calculates k-th order moment of truncated normal distribution.

Usage

```
truncatedNormalMoment(
  k = 1L,
  x_lower = numeric(0),
  x_upper = numeric(0),
  mean = 0,
  sd = 1,
  pdf_lower = numeric(0),
  cdf_lower = numeric(0),
  pdf_upper = numeric(0),
  cdf_upper = numeric(0),
  cdf_difference = numeric(0),
  return_all_moments = FALSE,
  is_validation = TRUE,
  is_parallel = FALSE,
  diff_type = "NO"
)
```

Arguments

k	non-negative integer moment order.
x_lower	numeric vector of lower truncation points.
x_upper	numeric vector of upper truncation points.
mean	numeric expected value.
sd	positive numeric standard deviation.
pdf_lower	non-negative numeric matrix of precalculated normal density functions with mean mean and standard deviation sd at points given by x_lower.
cdf_lower	non-negative numeric matrix of precalculated normal cumulative distribution functions with mean mean and standard deviation sd at points given by x_lower.
pdf_upper	non-negative numeric matrix of precalculated normal density functions with mean mean and standard deviation sd at points given by x_upper.
cdf_upper	non-negative numeric matrix of precalculated normal cumulative distribution functions with mean mean and standard deviation sd at points given by x_upper.
cdf_difference	non-negative numeric matrix of precalculated cdf_upper-cdf_lower values.

<code>return_all_moments</code>	logical; if TRUE, function returns the matrix of moments of normally distributed random variable with mean = mean and standard deviation = sd under lower and upper truncation points <code>x_lower</code> and <code>x_upper</code> correspondingly. Note that element in i-th row and j-th column of this matrix corresponds to the i-th observation (j-1)-th order moment.
<code>is_validation</code>	logical value indicating whether function input arguments should be validated. Set it to FALSE for slight performance boost (default value is TRUE).
<code>is_parallel</code>	if TRUE then multiple cores will be used for some calculations. It usually provides speed advantage for large enough samples (about more than 1000 observations).
<code>diff_type</code>	string indicating the type of the argument the moment should be differentiated respect to before return. Default value is "NO" so the moments itself will be returned. Alternative values are "mean", "sd", <code>x_lower</code> and <code>x_upper</code> .

Details

This function estimates k-th order moment of normal distribution which mean equals to mean and standard deviation equals to sd truncated at points given by `x_lower` and `x_upper`. Note that the function is vectorized so you can provide `x_lower` and `x_upper` as vectors of equal size. If vectors values for `x_lower` and `x_upper` are not provided then their default values will be set to $-(.Machine$double.xmin * 0.99)$ and $(.Machine$double.xmax * 0.99)$ correspondingly.

Note that parameter k value automatically converts to integer. So passing non-integer k value will not cause any errors but the calculations will be performed for rounded k value only.

If you have precalculated density or cumulative distribution functions at standardized truncation points (subtract mean and then divide by sd) then provide them through `pdf_lower`, `pdf_upper`, `cdf_lower` and `cdf_upper` arguments in order to decrease number of calculations.

Value

This function returns vector of k-th order moments for normally distributed random variable with mean = mean and standard deviation = sd under `x_lower` and `x_upper` truncation points `x_lower` and `x_upper` correspondingly. If `return_all_moments` is TRUE then see this argument description above for output details.

Examples

```
## Calculate 5-th order moment of three truncated normal random variables (x1,x2,x3)
## which mean is 5 and standard deviation is 3.
## These random variables truncation points are given as follows:-1<x1<1, 0<x2<2, 1<x3<3.
k <- 3
x_lower <- c(-1, 0, 1, -Inf, -Inf)
x_upper <- c(1, 2, 3, 2, Inf)
mean <- 3
sd <- 5

# get the moments
truncatedNormalMoment(k, x_lower, x_upper, mean, sd)
```

```
# get matrix of (0-5)-th moments (columns) for each variable (rows)
truncatedNormalMoment(k, x_lower, x_upper, mean, sd, return_all_moments = TRUE)

# get the moments derivatives respect to mean
truncatedNormalMoment(k, x_lower, x_upper, mean, sd, diff_type = "mean")

# get the moments derivatives respect to standard deviation
truncatedNormalMoment(k, x_lower, x_upper, mean, sd, diff_type = "sd")
```

Index

AIC.hpaBinary, [3](#), [23](#)
AIC.hpaML, [3](#), [28](#)
AIC.hpaSelection, [4](#), [35](#)
AIC_hpaBinary, [4](#)
AIC_hpaML, [5](#)
AIC_hpaSelection, [5](#)

dhp, [6](#), [25](#)
dhpDiff, [8](#), [9](#), [40](#)
dnorm_parallel, [11](#)
dtrhp, [12](#)

ehp, [15](#)
etrhp, [17](#)

ga, [21](#), [22](#), [26](#), [27](#), [32](#), [33](#)
gaControl, [22](#), [27](#), [33](#)
glm, [20](#)

hpaBinary, [19](#), [57–60](#), [66](#), [67](#)
hpaML, [24](#), [57–60](#), [66](#), [68](#)
hpaSelection, [30](#), [57–60](#), [67](#), [68](#)

ihp, [37](#)
ihpDiff, [9](#), [39](#), [40](#)
itrhp, [43](#)

logLik.hpaBinary, [23](#), [46](#)
logLik.hpaML, [28](#), [46](#)
logLik.hpaSelection, [35](#), [47](#)
logLik_hpaBinary, [47](#)
logLik_hpaML, [47](#)
logLik_hpaSelection, [48](#)

mecdf, [48](#)

normalMoment, [48](#)

optim, [21](#), [22](#), [26](#), [27](#), [32](#), [33](#)

php, [50](#)

plot.hpaBinary, [23](#), [52](#)
plot.hpaML, [53](#)
plot.hpaSelection, [35](#), [53](#)
plot_hpaBinary, [54](#)
plot_hpaSelection, [54](#)
pnorm_parallel, [55](#)
polynomialIndex, [55](#)
predict.hpaBinary, [23](#), [56](#)
predict.hpaML, [28](#), [57](#)
predict.hpaSelection, [35](#), [58](#)
predict_hpaBinary, [59](#)
predict_hpaML, [59](#)
predict_hpaSelection, [60](#)
print.hpaBinary, [61](#)
print.hpaML, [61](#)
print.hpaSelection, [62](#)
print.summary.hpaBinary, [62](#)
print.summary.hpaML, [63](#)
print.summary.hpaSelection, [63](#)
print_summary_hpaBinary, [64](#)
print_summary_hpaML, [65](#)
print_summary_hpaSelection, [65](#)
printPolynomial, [64](#)

summary.hpaBinary, [23](#), [66](#)
summary.hpaML, [28](#), [66](#)
summary.hpaSelection, [35](#), [67](#)
summary_hpaBinary, [67](#)
summary_hpaML, [68](#)
summary_hpaSelection, [68](#)

truncatedNormalMoment, [69](#)