

Package ‘gamCopula’

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Description Implementation of various inference and simulation tools to apply generalized additive models to bivariate dependence structures and non-simplified vine copulas.

License GPL-3

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R topics documented:

gamCopula-package	2
AIC.gamBiCop	8
BIC.gamBiCop	9
BiCopEta2Par	9

BiCopPar2Eta	10
condBiCopSim	11
dim.gamVine	13
EDF	14
formula.gamBiCop	15
gamBiCop	15
gamBiCop-class	16
gamBiCopCDF	16
gamBiCopFit	18
gamBiCopPDF	21
gamBiCopPredict	23
gamBiCopSelect	25
gamBiCopSimulate	28
gamVine	29
gamVine-class	30
gamVineCopSelect	31
gamVineFamily	35
gamVineNormalize	35
gamVineSeqFit	36
gamVineSimulate	39
gamVineStructureSelect	41
logLik.gamBiCop	44
nobs.gamBiCop	44
plot.gamBiCop	45
plot.gamVine	46
RVM2GVC	46
summary.gamBiCop	47
summary.gamVine	47
Index	49

gamCopula-package	<i>Generalized Additive Models for Bivariate Conditional Dependence Structures and Vine Copulas</i>
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Description

This package implements inference and simulation tools to apply generalized additive models to bivariate dependence structures and vine copulas.

More references in Vatter and Chavez-Demoulin (2015), and Vatter and Nagler (2016).

Details

Package:	gamCopula
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Author(s)

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References

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Vatter, T. and V. Chavez-Demoulin (2015) Generalized Additive Models for Conditional Dependence Structures. *Journal of Multivariate Analysis*, 141, 147-167.

Vatter, T. and T. Nagler (2016) Generalized additive models for non-simplified pair-copula constructions. <https://arxiv.org/abs/1608.01593>

Wood, S.N. (2004) Stable and efficient multiple smoothing parameter estimation for generalized additive models. *Journal of the American Statistical Association*, 99, 673-686.

Wood, S.N. (2006) *Generalized Additive Models: an introduction with R*. Chapman and Hall/CRC.

See Also

The present package is heavily relying on the [mgcv](#) and [VineCopula](#) packages, as it basically extends and mix both of them.

Examples

```
##### A gamBiCop example
require(copula)
require(mgcv)
set.seed(0)

## Simulation parameters (sample size, correlation between covariates,
## Gaussian copula family)
n <- 5e2
rho <- 0.5
fam <- 1

## A calibration surface depending on three variables
eta0 <- 1
calib.surf <- list(
```

```

calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
  Tm <- (Tf - Ti)/2
  a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
  return(a + b * (t - Tm)^2)},
calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
  a <- b * (1 - 2 * Tf * pi/(f * Tf * pi +
    cos(2 * f * pi * (Tf - Ti))
    - cos(2 * f * pi * Ti)))
  return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti))/2)},
calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
  Tm <- (Tf - Ti)/2
  a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
  return(a + b * exp(-(t - Tm)^2/(2 * s^2)))}

## Display the calibration surface
par(mfrow = c(1, 3), pty = "s", mar = c(1, 1, 4, 1))
u <- seq(0, 1, length.out = 100)
sel <- matrix(c(1, 1, 2, 2, 3, 3), ncol = 2)
jet.colors <- colorRamp(c("#00007F", "blue", "#007FFF", "cyan", "#7FFF7F",
  "yellow", "#FF7F00", "red", "#7F0000"))
jet <- function(x) rgb(jet.colors(exp(x/3)/(1 + exp(x/3))),
  maxColorValue = 255)
for (k in 1:3) {
  tmp <- outer(u, u, function(x, y)
    eta0 + calib.surf[[sel[k,1]]](x) + calib.surf[[sel[k, 2]]](y))
  persp(u, u, tmp, border = NA, theta = 60, phi = 30, zlab = "",
    col = matrix(jet(tmp), nrow = 100),
    xlab = paste("X", sel[k, 1], sep = ""),
    ylab = paste("X", sel[k,2], sep = ""),
    main = paste("eta0+f", sel[k, 1],
      "(X", sel[k, 1], ") +f", sel[k, 2],
      "(X", sel[k, 2], ")")", sep = ""))
}

## 3-dimensional matrix X of covariates
covariates.distr <- mvdc(normalCopula(rho, dim = 3),
  c("unif"), list(list(min = 0, max = 1)),
  marginsIdentical = TRUE)
X <- rMvdc(n, covariates.distr)

## U in [0,1]x[0,1] with copula parameter depending on X
U <- condBiCopSim(fam, function(x1,x2,x3) {eta0+sum(mapply(function(f,x)
  f(x), calib.surf, c(x1,x2,x3)))}, X[,1:3], par2 = 6, return.par = TRUE)

## Merge U and X
data <- data.frame(U$data,X)
names(data) <- c(paste("u",1:2,sep=""),paste("x",1:3,sep=""))

## Display the data
dev.off()
plot(data[, "u1"], data[, "u2"], xlab = "U1", ylab = "U2")

## Model fit with a basis size (arguably) too small

```

```

## and unpenalized cubic splines
pen <- FALSE
basis0 <- c(3, 4, 4)
formula <- ~s(x1, k = basis0[1], bs = "cr", fx = !pen) +
  s(x2, k = basis0[2], bs = "cr", fx = !pen) +
  s(x3, k = basis0[3], bs = "cr", fx = !pen)
system.time(fit0 <- gamBiCopFit(data, formula, fam))

## Model fit with a better basis size and penalized cubic splines (via min GCV)
pen <- TRUE
basis1 <- c(3, 10, 10)
formula <- ~s(x1, k = basis1[1], bs = "cr", fx = !pen) +
  s(x2, k = basis1[2], bs = "cr", fx = !pen) +
  s(x3, k = basis1[3], bs = "cr", fx = !pen)
system.time(fit1 <- gamBiCopFit(data, formula, fam))

## Extract the gamBiCop objects and show various methods
(res <- sapply(list(fit0,fit1), function(fit){fit$res}))
metds <- list('logLik'=logLik,'AIC'=AIC,'BIC'=BIC,'EDF'=EDF)
lapply(res, function(x) sapply(metds, function(f) f(x)))

## Comparison between fitted, true smooth and spline approximation for each
## true smooth function for the two basis sizes
fitted <- lapply(res, function(x) gamBiCopPredict(x, data.frame(x1=u,x2=u,x3=u),
  type = "terms")$calib)

true <- vector("list", 3)
for (i in 1:3) {
  y <- eta0+calib.surf[[i]](u)
  true[[i]]$true <- y - eta0
  temp <- gam(y ~ s(u, k = basis0[i], bs = "cr", fx = TRUE))
  true[[i]]$approx <- predict.gam(temp, type = "terms")
  temp <- gam(y ~s(u, k = basis1[i], bs = "cr", fx = FALSE))
  true[[i]]$approx2 <- predict.gam(temp, type = "terms")
}

## Display results
par(mfrow = c(1, 3), pty = "s")
yy <- range(true, fitted)
yy[1] <- yy[1] * 1.5
for(k in 1:3){
  plot(u, true[[k]]$true, type = "l", ylim = yy,
    xlab = paste("Covariate",k), ylab = paste("Smooth",k))
  lines(u, true[[k]]$approx, col = "red", lty = 2)
  lines(u, fitted[[1]][, k], col = "red")
  lines(u, fitted[[2]][, k], col = "green")
  lines(u, true[[k]]$approx2, col = "green", lty = 2)
  legend("bottomleft", cex = 0.6, lty = c(1, 1, 2, 1, 2),
    c("True", "Fitted", "Appox 1", "Fitted 2", "Approx 2"),
    col = c("black", "red", "red", "green", "green"))
}

##### A gamVine example

```

```

set.seed(0)

## Simulation parameters
# Sample size
n <- 1e3
# Copula families
familyset <- c(1:2,301:304,401:404)
# Define a 4-dimensional R-vine tree structure matrix
d <- 4
Matrix <- c(2,3,4,1,0,3,4,1,0,0,4,1,0,0,0,1)
Matrix <- matrix(Matrix,d,d)
nnames <- paste("X", 1:d, sep = "")

## A function factory
eta0 <- 1
calib.surf <- list(
  calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
    Tm <- (Tf - Ti)/2
    a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
    return(a + b * (t - Tm)^2)},
  calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
    a <- b * (1 - 2 * Tf * pi/(f * Tf * pi +
      cos(2 * f * pi * (Tf - Ti))
      - cos(2 * f * pi * Ti)))
    return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti))/2)},
  calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
    Tm <- (Tf - Ti)/2
    a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
    return(a + b * exp(-(t - Tm)^2/(2 * s^2)))})

## Create the model
# Define gam-vine model list
count <- 1
model <- vector(mode = "list", length = d*(d-1)/2)
sel <- seq(d,d^2-d, by = d)

# First tree
for (i in 1:(d-1)) {
  # Select a copula family
  family <- sample(familyset, 1)
  model[[count]]$family <- family

  # Use the canonical link and a randomly generated parameter
  if (is.element(family,c(1,2))) {
    model[[count]]$par <- tanh(rnorm(1)/2)
    if (family == 2) {
      model[[count]]$par2 <- 2+exp(rnorm(1))
    }
  } else {
    if (is.element(family,c(401:404))) {
      rr <- rnorm(1)
      model[[count]]$par <- sign(rr)*(1+abs(rr))
    } else {

```

```

    model[[count]]$par <- rnorm(1)
  }
  model[[count]]$par2 <- 0
}
count <- count + 1
}

# A dummy dataset
data <- data.frame(u1 = runif(1e2), u2 = runif(1e2), matrix(runif(1e2*d),1e2,d))

# Trees 2 to (d-1)
for(j in 2:(d-1)){
  for(i in 1:(d-j)){
    # Select a copula family
    family <- sample(familyset, 1)

    # Select the conditioning set and create a model formula
    cond <- nnames[sort(Matrix[(d-j+2):d,i])]
    tmpform <- paste("~",paste(paste("s(", cond, ", k=10, bs='cr')",
                                   sep = ""), collapse=" + "))

    l <- length(cond)
    temp <- sample(3, l, replace = TRUE)

    # Spline approximation of the true function
    m <- 1e2
    x <- matrix(seq(0,1,length.out=m), nrow = m, ncol = 1)
    if(l != 1){
      tmp.fct <- paste("function(x){eta0+",
                      paste(sapply(1:l, function(x)
                                   paste("calib.surf[[",temp[x],"]](x[,x,""],
                                       sep="")), collapse="+"),")",sep="")
      tmp.fct <- eval(parse(text = tmp.fct))
      x <- eval(parse(text = paste0("expand.grid(",
                                   paste0(rep("x",l), collapse = ","),")",
                                   collapse = "")))

      y <- apply(x,1,tmp.fct)
    }else{
      tmp.fct <- function(x) eta0+calib.surf[[temp]](x)
      colnames(x) <- cond
      y <- tmp.fct(x)
    }

    # Estimate the gam model
    form <- as.formula(paste0("y", tmpform))
    dd <- data.frame(y, x)
    names(dd) <- c("y", cond)
    b <- gam(form, data = dd)
    #plot(x[,1],(y-fitted(b))/y)

    # Create a dummy gamBiCop object
    tmp <- gamBiCopFit(data = data, formula = form, family = 1, n.iters = 1)$res

    # Update the copula family and the model coefficients

```

```

    attr(tmp, "model")$coefficients <- coefficients(b)
    attr(tmp, "model")$smooth <- b$smooth
    attr(tmp, "family") <- family
    if (family == 2) {
      attr(tmp, "par2") <- 2+exp(rnorm(1))
    }
    model[[count]] <- tmp
    count <- count+1
  }
}

# Create the gamVineCopula object
GVC <- gamVine(Matrix=Matrix,model = model,names=nnames)
print(GVC)

## Not run:
## Simulate and fit the model
sim <- gamVineSimulate(n, GVC)
fitGVC <- gamVineSeqFit(sim, GVC, verbose = TRUE)
fitGVC2 <- gamVineCopSelect(sim, Matrix, verbose = TRUE)

## Plot the results
par(mfrow=c(3,4))
plot(GVC, ylim = c(-2.5,2.5))

plot(fitGVC, ylim = c(-2.5,2.5))

plot(fitGVC2, ylim = c(-2.5,2.5))
## End(Not run)

```

AIC.gamBiCop

Akaike's An Information Criterion for a gamBiCop Object

Description

Function calculating Akaike's 'An Information Criterion' (AIC) for an object of the class `gamBiCop` (note that the models are usually fitted by penalized likelihood maximization).

Usage

```
## S4 method for signature 'gamBiCop'
AIC(object, ..., k = 2)
```

Arguments

<code>object</code>	An object of the class <code>gamBiCop</code> .
<code>...</code>	un-used in this class
<code>k</code>	numeric, the penalty per parameter to be used; the default <code>k = 2</code> is the classical AIC.

Value

A numeric value with the corresponding AIC.

See Also

[AIC](#) and [BIC](#).

 BIC.gamBiCop

Schwarz's Bayesian Information Criterion for a gamBiCop Object

Description

Function calculating the Schwarz's Bayesian Information Criterion (BIC) for an object of the class [gamBiCop](#) (note that the models are usually fitted by penalized likelihood maximization).

Usage

```
## S4 method for signature 'gamBiCop'
BIC(object, ...)
```

Arguments

object	An object of the class gamBiCop .
...	un-used in this class

Value

A numeric value with the corresponding BIC.

See Also

[AIC](#) and [BIC](#).

 BiCopEta2Par

Copula Parameter of a Bivariate Copula for a Given Value of the Calibration Function

Description

Computes the (first) copula parameter of a bivariate copula for a given value of the calibration function (eta).

Usage

```
BiCopEta2Par(family, eta)
```

Arguments

family	A copula family: 1 Gaussian, 2 Student t, 301 Double Clayton type I (standard and rotated 90 degrees), 302 Double Clayton type II (standard and rotated 270 degrees), 303 Double Clayton type III (survival and rotated 90 degrees), 304 Double Clayton type IV (survival and rotated 270 degrees), 401 Double Gumbel type I (standard and rotated 90 degrees), 402 Double Gumbel type II (standard and rotated 270 degrees), 403 Double Gumbel type III (survival and rotated 90 degrees), 404 Double Gumbel type IV (survival and rotated 270 degrees).
eta	The calibration function.

Value

The value of the first copula parameter, depending on the copula parameter and family as:

- 1 Gaussian, $f(x) = \tanh(x/2)$
- 2 Student t, $f(x) = \tanh(x/2)$
- 301 Double Clayton type I (standard and rotated 90 degrees), $f(x) = x$
- 302 Double Clayton type II (standard and rotated 270 degrees), $f(x) = x$
- 303 Double Clayton type III (survival and rotated 90 degrees), $f(x) = x$
- 304 Double Clayton type IV (survival and rotated 270 degrees), $f(x) = x$
- 401 Double Gumbel type I (standard and rotated 90 degrees), $f(x) = x*(1+abs(x))/abs(x)$
- 402 Double Gumbel type II (standard and rotated 270 degrees), $f(x) = x*(1+abs(x))/abs(x)$
- 403 Double Gumbel type III (survival and rotated 90 degrees), $f(x) = x*(1+abs(x))/abs(x)$
- 404 Double Gumbel type IV (survival and rotated 270 degrees) $f(x) = x*(1+abs(x))/abs(x)$.

See Also

[BiCopEta2Par](#) or [BiCopPar2Tau](#) and [BiCopTau2Par](#) from [VineCopula](#).

BiCopPar2Eta

Calibration Function of a Bivariate Copula for a Given Parameter's Value

Description

Computes the calibration function (eta) of a bivariate copula for a given value of the (first) copula parameter.

Usage

BiCopPar2Eta(family, par)

Arguments

family	A copula family: 1 Gaussian, 2 Student t, 301 Double Clayton type I (standard and rotated 90 degrees), 302 Double Clayton type II (standard and rotated 270 degrees), 303 Double Clayton type III (survival and rotated 90 degrees), 304 Double Clayton type IV (survival and rotated 270 degrees), 401 Double Gumbel type I (standard and rotated 90 degrees), 402 Double Gumbel type II (standard and rotated 270 degrees), 403 Double Gumbel type III (survival and rotated 90 degrees), 404 Double Gumbel type IV (survival and rotated 270 degrees).
par	The (first) copula parameter

Value

The value of the calibration function, depending on the copula parameter and family as:

- 1 Gaussian, $f(x) = 2 \cdot \operatorname{atanh}(x)$
- 2 Student t, $f(x) = 2 \cdot \operatorname{atanh}(x)$
- 301 Double Clayton type I (standard and rotated 90 degrees), $f(x) = x$
- 302 Double Clayton type II (standard and rotated 270 degrees), $f(x) = x$
- 303 Double Clayton type III (survival and rotated 90 degrees), $f(x) = x$
- 304 Double Clayton type IV (survival and rotated 270 degrees), $f(x) = x$
- 401 Double Gumbel type I (standard and rotated 90 degrees), $f(x) = x \cdot (1 - 1/\operatorname{abs}(x))$
- 402 Double Gumbel type II (standard and rotated 270 degrees), $f(x) = x \cdot (1 - 1/\operatorname{abs}(x))$
- 403 Double Gumbel type III (survival and rotated 90 degrees), $f(x) = x \cdot (1 - 1/\operatorname{abs}(x))$
- 404 Double Gumbel type IV (survival and rotated 270 degrees) $f(x) = x \cdot (1 - 1/\operatorname{abs}(x))$.

See Also

[BiCopEta2Par](#) or [BiCopPar2Tau](#) and [BiCopTau2Par](#) from [VineCopula](#).

 condBiCopSim

Simulation from a Conditional Bivariate Copula

Description

Simulates from a conditional bivariate copula, where each copula parameter takes a different value, depending on the calibration function and covariates.

Usage

```
condBiCopSim(family, calib.fnc, X, par2 = 0, return.par = TRUE,
             tau = TRUE)
```

Arguments

family	family A copula family: 1 Gaussian, 2 Student t, 3 Clayton, 4 Gumbel, 5 Frank, 13 Survival Clayton, 14 Survival Gumbel, 23 Rotated (90 degrees) Clayton, 24 Rotated (90 degrees) Gumbel, 33 Rotated (270 degrees) Clayton and 34 Rotated (270 degrees) Gumbel.
calib.fnc	A calibration function.
X	A vector (if calib.fnc takes a single argument) or matrix (if calib.fnc takes multiple arguments) of covariates values.
par2	The second copula parameter (for the Student t), default par2 = 0.
return.par	Should the parameter (and calibration function) be returned as well (default return.par = TRUE)?
tau	Should the calibration function (and the model) be specified for the copula parameter or Kendall's tau (default tau = TRUE)?

Value

If return.par = TRUE, then the function returns a list with:

- data, a matrix with two columns containing the simulated data,
- par, a vector containing the values of the copula parameter,
- and eta, a vector containing the values of the calibration function.

If return.par = FALSE, then the function simply returns data, a matrix with two columns containing the simulated data.

See Also

[gamBiCopFit](#) and [gamBiCopSimulate](#).

Examples

```
require(copula)
set.seed(0)

## Simulation parameters (sample size, correlation between covariates,
## Gaussian copula family)
n <- 2e2
rho <- 0.5
fam <- 1

## A calibration surface depending on three variables
eta0 <- 1
calib.surf <- list(
  calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
    Tm <- (Tf - Ti)/2
    a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
    return(a + b * (t - Tm)^2)},
  calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
```

```

a <- b * (1 - 2 * Tf * pi / (f * Tf * pi +
                           cos(2 * f * pi * (Tf - Ti))
                           - cos(2 * f * pi * Ti)))
return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti))/2)},
calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
  Tm <- (Tf - Ti)/2
  a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
  return(a + b * exp(-(t - Tm)^2/(2 * s^2))))}

## Display the calibration surface
par(mfrow = c(1, 3), pty = "s", mar = c(1, 1, 4, 1))
u <- seq(0, 1, length.out = 100)
sel <- matrix(c(1, 1, 2, 2, 3, 3), ncol = 2)
jet.colors <- colorRamp(c("#00007F", "blue", "#007FFF", "cyan", "#7FFF7F",
                          "yellow", "#FF7F00", "red", "#7F0000"))
jet <- function(x) rgb(jet.colors(exp(x/3)/(1 + exp(x/3))),
                       maxColorValue = 255)

for (k in 1:3) {
  tmp <- outer(u, u, function(x, y)
               eta0 + calib.surf[[sel[k,1]]](x) + calib.surf[[sel[k, 2]]](y))
  persp(u, u, tmp, border = NA, theta = 60, phi = 30, zlab = "",
        col = matrix(jet(tmp), nrow = 100),
        xlab = paste("X", sel[k, 1], sep = ""),
        ylab = paste("X", sel[k,2], sep = ""),
        main = paste("eta0+f", sel[k, 1],
                     "(X", sel[k, 1], ") +f", sel[k, 2],
                     "(X", sel[k, 2], ") ", sep = ""))
}

## 3-dimensional matrix X of covariates
covariates.distr <- mvdc(normalCopula(rho, dim = 3),
                        c("unif"), list(list(min = 0, max = 1)),
                        marginsIdentical = TRUE)
X <- rMvdc(n, covariates.distr)

## U in [0,1]x[0,1] with copula parameter depending on X
U <- condBiCopSim(fam, function(x1,x2,x3) {eta0+sum(mapply(function(f,x)
  f(x), calib.surf, c(x1,x2,x3)))}, X[,1:3], par2 = 6, return.par = TRUE)

## Merge U and X
data <- data.frame(U$data,X)
names(data) <- c(paste("u",1:2,sep=""),paste("x",1:3,sep=""))

## Display the data
dev.off()
plot(data[, "u1"], data[, "u2"], xlab = "U1", ylab = "U2")

```

Description

Retrieve the dimension of an object of the class `gamVine`.

Usage

```
## S4 method for signature 'gamVine'  
dim(x)
```

Arguments

x An object of the class `gamVine`.

Value

Dimension of the `gamVine` object.

See Also

`gamVine`.

EDF

Equivalent Degrees of Freedom for an Object of the Class `gamBiCop`

Description

Function calculating the Equivalent Degrees of Freedom (EDF) for a `gamBiCop` object. It basically sums the edf of the `gamObject` for each smooth component.

Usage

```
EDF(object)
```

Arguments

object An object of the class `gamBiCop`.

Value

Estimated degrees of freedom for each smooth component.

formula.gamBiCop	<i>Model Formula of the gamBiCop Object</i>
------------------	---

Description

Extracts the `gam` formula from an object of the class `gamBiCop`. This function is a wrapper to `formula.gam` from the `mgcv` package.

Usage

```
## S4 method for signature 'gamBiCop'
formula(x, ...)
```

Arguments

<code>x</code>	An object of the class <code>gamBiCop</code> .
<code>...</code>	un-used in this class

See Also

`formula.gam` function from the `mgcv` package.

gamBiCop	<i>Construction of a gamBiCop Class Object</i>
----------	--

Description

Constructs an object of the class `gamBiCop`.

Usage

```
gamBiCop(family, model, par2 = 0, tau = TRUE)
```

Arguments

<code>family</code>	A copula family: 1 Gaussian, 2 Student t, 5 Frank, 301 Double Clayton type I (standard and rotated 90 degrees), 302 Double Clayton type II (standard and rotated 270 degrees), 303 Double Clayton type III (survival and rotated 90 degrees), 304 Double Clayton type IV (survival and rotated 270 degrees), 401 Double Gumbel type I (standard and rotated 90 degrees), 402 Double Gumbel type II (standard and rotated 270 degrees), 403 Double Gumbel type III (survival and rotated 90 degrees), 404 Double Gumbel type IV (survival and rotated 270 degrees).
<code>model</code>	A <code>gamObject</code> as return by the <code>gam</code> function from the <code>mgcv</code> package.
<code>par2</code>	Second parameter for the Student t-copula.
<code>tau</code>	FALSE for a calibration function specified for the Copula parameter or TRUE (default) for a calibration function specified for Kendall's tau.

Value

An object of the class [gamBiCop](#).

See Also

[gamBiCop](#), [gamBiCopFit](#), [gamBiCopPredict](#) and [gamBiCopSimulate](#).

gamBiCop-class	<i>The gamBiCop Class</i>
----------------	---------------------------

Description

`gamBiCop` is an S4 class to store a Generalized Additive Model for bivariate copula a parameter or Kendall's tau. Objects can be created by calls of the form `new("gamBiCop", ...)`, or by function [gamBiCop](#).

Slots

`family` A copula family: 1 Gaussian, 2 Student t, 5 Frank, 301 Double Clayton type I (standard and rotated 90 degrees), 302 Double Clayton type II (standard and rotated 270 degrees), 303 Double Clayton type III (survival and rotated 90 degrees), 304 Double Clayton type IV (survival and rotated 270 degrees), 401 Double Gumbel type I (standard and rotated 90 degrees), 402 Double Gumbel type II (standard and rotated 270 degrees), 403 Double Gumbel type III (survival and rotated 90 degrees), 404 Double Gumbel type IV (survival and rotated 270 degrees).

`model` A [gamObject](#) as return by the `gam` function from the `mgcv` package.

`par2` Second parameter for the Student t-copula.

`tau` FALSE (default) for a calibration function specified for the Copula parameter or TRUE for a calibration function specified for Kendall's tau.

See Also

[gamBiCopFit](#), [gamBiCopPredict](#) and [gamBiCopSimulate](#).

gamBiCopCDF	<i>Conditional distribution function of a Generalized Additive model for the copula parameter or Kendall's tau</i>
-------------	--

Description

This function returns the distribution function of a bivariate conditional copula, where either the copula parameter or the Kendall's tau is modeled as a function of the covariates.

Usage

```
gamBiCopCDF(object, newdata = NULL)
```

Arguments

object [gamBiCop-class](#) object.

newdata (Same as in [predict.gam](#) from the [mgcv](#) package) A matrix or data frame containing the values of the model covariates at which predictions are required. If this is not provided then the distribution corresponding to the original data are returned. If `newdata` is provided then it should contain all the variables needed for prediction: a warning is generated if not.

Value

The conditional density.

See Also

[gamBiCop](#) and [gamBiCopPredict](#).

Examples

```
require(copula)
set.seed(0)

## Simulation parameters (sample size, correlation between covariates,
## Gaussian copula family)
n <- 2e2
rho <- 0.5
fam <- 1

## A calibration surface depending on three variables
eta0 <- 1
calib.surf <- list(
  calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
    Tm <- (Tf - Ti)/2
    a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
    return(a + b * (t - Tm)^2)},
  calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
    a <- b * (1 - 2 * Tf * pi / (f * Tf * pi +
      cos(2 * f * pi * (Tf - Ti))
      - cos(2 * f * pi * Ti)))
    return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti))/2)},
  calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
    Tm <- (Tf - Ti)/2
    a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
    return(a + b * exp(-(t - Tm)^2/(2 * s^2)))})

## 3-dimensional matrix X of covariates
covariates.distr <- mvdc(normalCopula(rho, dim = 3),
  c("unif"), list(list(min = 0, max = 1)),
```

```

                                marginsIdentical = TRUE)
X <- rMvdc(n, covariates.distr)
colnames(X) <- paste("x",1:3,sep="")

## U in [0,1]x[0,1] with copula parameter depending on X
U <- condBiCopSim(fam, function(x1,x2,x3) {eta0+sum(mapply(function(f,x)
  f(x), calib.surf, c(x1,x2,x3)))}, X[,1:3], par2 = 6, return.par = TRUE)

## Merge U and X
data <- data.frame(U$data,X)
names(data) <- c(paste("u",1:2,sep=""),paste("x",1:3,sep=""))

## Model fit with penalized cubic splines (via min GCV)
basis <- c(3, 10, 10)
formula <- ~s(x1, k = basis[1], bs = "cr") +
  s(x2, k = basis[2], bs = "cr") +
  s(x3, k = basis[3], bs = "cr")
system.time(fit <- gamBiCopFit(data, formula, fam))

## Evaluate the conditional density
gamBiCopCDF(fit$res)

```

gamBiCopFit

Maximum penalized likelihood estimation of a Generalized Additive model for the copula parameter or Kendall's tau.

Description

This function estimates the parameter(s) of a Generalized Additive model (gam) for the copula parameter or Kendall's tau. It solves the maximum penalized likelihood estimation for the copula families supported in this package by reformulating each Newton-Raphson iteration as a generalized ridge regression, which is solved using the [mgcv](#) package.

Usage

```
gamBiCopFit(data, formula = ~1, family = 1, tau = TRUE, method = "FS",
  tol.rel = 0.001, n.iters = 10, verbose = FALSE, ...)
```

Arguments

data	A list, data frame or matrix containing the model responses, (u1,u2) in [0,1]x[0,1], and covariates required by the formula.
formula	A gam formula (see gam , formula.gam and gam.models from mgcv).
family	A copula family: 1 Gaussian, 2 Student t, 5 Frank, 301 Double Clayton type I (standard and rotated 90 degrees), 302 Double Clayton type II (standard and rotated 270 degrees), 303 Double Clayton type III (survival and rotated 90 degrees), 304 Double Clayton type IV (survival and rotated 270 degrees), 401

	Double Gumbel type I (standard and rotated 90 degrees), 402 Double Gumbel type II (standard and rotated 270 degrees), 403 Double Gumbel type III (survival and rotated 90 degrees), 404 Double Gumbel type IV (survival and rotated 270 degrees).
tau	FALSE (default) for a calibration function specified for the Copula parameter or TRUE for a calibration function specified for Kendall's tau.
method	'NR' for Newton-Raphson and 'FS' for Fisher-scoring (default).
tol.rel	Relative tolerance for 'FS'/'NR' algorithm.
n.iters	Maximal number of iterations for 'FS'/'NR' algorithm.
verbose	TRUE if informations should be printed during the estimation and FALSE (default) for a silent version.
...	Additional parameters to be passed to <code>gam</code> from <code>mgcv</code> .

Value

`gamBiCopFit` returns a list consisting of

res	S4 <code>gamBiCop-class</code> object.
method	'FS' for Fisher-scoring (default) and 'NR' for Newton-Raphson.
tol.rel	relative tolerance for 'FS'/'NR' algorithm.
n.iters	maximal number of iterations for 'FS'/'NR' algorithm.
trace	the estimation procedure's trace.
conv	0 if the algorithm converged and 1 otherwise.

See Also

[gamBiCop](#) and [gamBiCopSimulate](#).

Examples

```
require(copula)
require(mgcv)
set.seed(0)

## Simulation parameters (sample size, correlation between covariates,
## Gaussian copula family)
n <- 5e2
rho <- 0.5
fam <- 1

## A calibration surface depending on three variables
eta0 <- 1
calib.surf <- list(
  calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
    Tm <- (Tf - Ti)/2
    a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
    return(a + b * (t - Tm)^2)},
```

```

calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
  a <- b * (1 - 2 * Tf * pi / (f * Tf * pi +
    cos(2 * f * pi * (Tf - Ti))
    - cos(2 * f * pi * Ti)))
  return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti)/2)},
calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
  Tm <- (Tf - Ti)/2
  a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
  return(a + b * exp(-(t - Tm)^2/(2 * s^2))))

## Display the calibration surface
par(mfrow = c(1, 3), pty = "s", mar = c(1, 1, 4, 1))
u <- seq(0, 1, length.out = 100)
sel <- matrix(c(1, 1, 2, 2, 3, 3), ncol = 2)
jet.colors <- colorRamp(c("#00007F", "blue", "#007FFF", "cyan", "#7FFF7F",
  "yellow", "#FF7F00", "red", "#7F0000"))
jet <- function(x) rgb(jet.colors(exp(x/3)/(1 + exp(x/3))),
  maxColorValue = 255)

for (k in 1:3) {
  tmp <- outer(u, u, function(x, y)
    eta0 + calib.surf[[sel[k,1]]](x) + calib.surf[[sel[k, 2]]](y))
  persp(u, u, tmp, border = NA, theta = 60, phi = 30, zlab = "",
    col = matrix(jet(tmp), nrow = 100),
    xlab = paste("X", sel[k, 1], sep = ""),
    ylab = paste("X", sel[k,2], sep = ""),
    main = paste("eta0+f", sel[k, 1],
      "(X", sel[k, 1], ") +f", sel[k, 2],
      "(X", sel[k, 2], ") ", sep = ""))
}

## 3-dimensional matrix X of covariates
covariates.distr <- mvdc(normalCopula(rho, dim = 3),
  c("unif"), list(list(min = 0, max = 1)),
  marginsIdentical = TRUE)
X <- rMvdc(n, covariates.distr)

## U in [0,1]x[0,1] with copula parameter depending on X
U <- condBiCopSim(fam, function(x1,x2,x3) {eta0+sum(mapply(function(f,x)
  f(x), calib.surf, c(x1,x2,x3)))}, X[,1:3], par2 = 6, return.par = TRUE)

## Merge U and X
data <- data.frame(U$data,X)
names(data) <- c(paste("u",1:2,sep=""),paste("x",1:3,sep=""))

## Display the data
dev.off()
plot(data[, "u1"], data[, "u2"], xlab = "U1", ylab = "U2")

## Model fit with a basis size (arguably) too small
## and unpenalized cubic spines
pen <- FALSE
basis0 <- c(3, 4, 4)
formula <- ~s(x1, k = basis0[1], bs = "cr", fx = !pen) +

```

```

s(x2, k = basis0[2], bs = "cr", fx = !pen) +
s(x3, k = basis0[3], bs = "cr", fx = !pen)
system.time(fit0 <- gamBiCopFit(data, formula, fam))

## Model fit with a better basis size and penalized cubic splines (via min GCV)
pen <- TRUE
basis1 <- c(3, 10, 10)
formula <- ~s(x1, k = basis1[1], bs = "cr", fx = !pen) +
  s(x2, k = basis1[2], bs = "cr", fx = !pen) +
  s(x3, k = basis1[3], bs = "cr", fx = !pen)
system.time(fit1 <- gamBiCopFit(data, formula, fam))

## Extract the gamBiCop objects and show various methods
(res <- sapply(list(fit0,fit1), function(fit){fit$res}))
metds <- list('logLik'=logLik, 'AIC'=AIC, 'BIC'=BIC, 'EDF'=EDF)
lapply(res, function(x) sapply(metds, function(f) f(x)))

## Comparison between fitted, true smooth and spline approximation for each
## true smooth function for the two basis sizes
fitted <- lapply(res, function(x) gamBiCopPredict(x, data.frame(x1=u,x2=u,x3=u),
  type = "terms")$calib)

true <- vector("list", 3)
for (i in 1:3) {
  y <- eta0+calib.surf[[i]](u)
  true[[i]]$true <- y - eta0
  temp <- gam(y ~ s(u, k = basis0[i], bs = "cr", fx = TRUE))
  true[[i]]$approx <- predict.gam(temp, type = "terms")
  temp <- gam(y ~s(u, k = basis1[i], bs = "cr", fx = FALSE))
  true[[i]]$approx2 <- predict.gam(temp, type = "terms")
}

## Display results
par(mfrow = c(1, 3), pty = "s")
yy <- range(true, fitted)
yy[1] <- yy[1] * 1.5
for(k in 1:3){
  plot(u, true[[k]]$true, type = "l", ylim = yy,
    xlab = paste("Covariate",k), ylab = paste("Smooth",k))
  lines(u, true[[k]]$approx, col = "red", lty = 2)
  lines(u, fitted[[1]][, k], col = "red")
  lines(u, fitted[[2]][, k], col = "green")
  lines(u, true[[k]]$approx2, col = "green", lty = 2)
  legend("bottomleft", cex = 0.6, lty = c(1, 1, 2, 1, 2),
    c("True", "Fitted", "Appox 1", "Fitted 2", "Appox 2"),
    col = c("black", "red", "red", "green", "green"))
}

```

Description

This function returns the density of a bivariate conditional copula, where either the copula parameter or the Kendall's tau is modeled as a function of the covariates.

Usage

```
gamBiCopPDF(object, newdata = NULL)
```

Arguments

object	gamBiCop-class object.
newdata	(Same as in predict.gam from the mgcv package) A matrix or data frame containing the values of the model covariates at which predictions are required. If this is not provided then the density corresponding to the original data are returned. If newdata is provided then it should contain all the variables needed for prediction: a warning is generated if not.

Value

The conditional density.

See Also

[gamBiCop](#) and [gamBiCopPredict](#).

Examples

```
require(copula)
set.seed(0)

## Simulation parameters (sample size, correlation between covariates,
## Gaussian copula family)
n <- 2e2
rho <- 0.5
fam <- 1

## A calibration surface depending on three variables
eta0 <- 1
calib.surf <- list(
  calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
    Tm <- (Tf - Ti)/2
    a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
    return(a + b * (t - Tm)^2)},
  calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
    a <- b * (1 - 2 * Tf * pi/(f * Tf * pi +
      cos(2 * f * pi * (Tf - Ti))
      - cos(2 * f * pi * Ti)))
    return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti))/2)},
  calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
    Tm <- (Tf - Ti)/2
    a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
```

```

    return(a + b * exp(-(t - Tm)^2/(2 * s^2)))})

## 3-dimensional matrix X of covariates
covariates.distr <- mvdc(normalCopula(rho, dim = 3),
                        c("unif"), list(list(min = 0, max = 1)),
                        marginsIdentical = TRUE)
X <- rMvdc(n, covariates.distr)
colnames(X) <- paste("x",1:3,sep="")

## U in [0,1]x[0,1] with copula parameter depending on X
U <- condBiCopSim(fam, function(x1,x2,x3) {eta0+sum(mapply(function(f,x)
  f(x), calib.surf, c(x1,x2,x3)))}, X[,1:3], par2 = 6, return.par = TRUE)

## Merge U and X
data <- data.frame(U$data,X)
names(data) <- c(paste("u",1:2,sep=""),paste("x",1:3,sep=""))

## Model fit with penalized cubic splines (via min GCV)
basis <- c(3, 10, 10)
formula <- ~s(x1, k = basis[1], bs = "cr") +
  s(x2, k = basis[2], bs = "cr") +
  s(x3, k = basis[3], bs = "cr")
system.time(fit <- gamBiCopFit(data, formula, fam))

## Evaluate the conditional density
gamBiCopPDF(fit$res)

```

gamBiCopPredict	<i>Predict method of a Generalized Additive model for the copula parameter or Kendall's tau</i>
-----------------	---

Description

Predict method of a Generalized Additive model for the copula parameter or Kendall's tau

Usage

```
gamBiCopPredict(object, newdata = NULL, target = "calib", alpha = 0,
  type = "link")
```

Arguments

object	gamBiCop-class object.
newdata	(Same as in predict.gam from the mgcv package) A matrix or data frame containing the values of the model covariates at which predictions are required. If this is not provided then predictions corresponding to the original data are returned. If newdata is provided then it should contain all the variables needed for prediction: a warning is generated if not.

target	Either 'calib', 'par' or 'tau' or a combination of those. 'calib' (default) corresponds to the calibration function, 'par' to the copula parameter and 'tau' to Kendall's tau.
alpha	In (0,1) to return the corresponding confidence interval.
type	(Similar as in predict.gam from the mgcv package, only active for type = 'calib'). When this has the value 'link' (default), the calibration function is returned. When type = 'terms' each component of the linear predictor is returned separately (possibly with standard errors): this includes parametric model components, followed by each smooth component, but excludes any offset and any intercept. When type = 'lpmatrix' then a matrix is returned which yields the values of the linear predictor (minus any offset) when post-multiplied by the parameter vector (in this case alpha is ignored).

Value

If target = 'calib', then a list with 1 item calib. If target = 'par', target = 'tau' or target = c('par', 'tau'), then a list with 2, 2 or 3 items, namely calib and par, tau and par, or calib, tau and par.

If alpha is in (0,1), then a additional items of the list are calib.CI as well as e.g. par.CI and/or tau.CI depending on the value of target.

Otherwise, if type = 'lpmatrix' (only active for type = 'calib'), then a matrix is returned which will give a vector of linear predictor values (minus any offset) at the supplied covariate values, when applied to the model coefficient vector (similar as [predict.gam](#) from the [mgcv](#)).

See Also

[gamBiCop](#) and [gamBiCopFit](#).

Examples

```
require(copula)
set.seed(0)

## Simulation parameters (sample size, correlation between covariates,
## Clayton copula family)
n <- 5e2
rho <- 0.5
fam <- 1

## A calibration surface depending on three variables
eta0 <- 1
calib.surf <- list(
  calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
    Tm <- (Tf - Ti)/2
    a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
    return(a + b * (t - Tm)^2)},
  calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
    a <- b * (1 - 2 * Tf * pi/(f * Tf * pi +
      cos(2 * f * pi * (Tf - Ti))
      - cos(2 * f * pi * Ti)))
```



```

    return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti))/2)},
  calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
    Tm <- (Tf - Ti)/2
    a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
    return(a + b * exp(-(t - Tm)^2/(2 * s^2)))})

## 3-dimensional matrix X of covariates
covariates.distr <- mvdc(normalCopula(rho, dim = 3),
                        c("unif"), list(list(min = 0, max = 1)),
                        marginsIdentical = TRUE)
X <- rMvdc(n, covariates.distr)
colnames(X) <- paste("x",1:3,sep="")

## U in [0,1]x[0,1] with copula parameter depending on X
U <- condBiCopSim(fam, function(x1,x2,x3) {eta0+sum(mapply(function(f,x)
  f(x), calib.surf, c(x1,x2,x3)))}, X[,1:3], par2 = 6, return.par = TRUE)

## Merge U and X
data <- data.frame(U$data,X)
names(data) <- c(paste("u",1:2,sep=""),paste("x",1:3,sep=""))

## Model fit with penalized cubic splines (via min GCV)
basis <- c(3, 10, 10)
formula <- ~s(x1, k = basis[1], bs = "cr") +
  s(x2, k = basis[2], bs = "cr") +
  s(x3, k = basis[3], bs = "cr")
system.time(fit <- gamBiCopFit(data, formula, fam))

## Extract the gamBiCop objects and show various methods
(res <- fit$res)
EDF(res)
pred <- gamBiCopPredict(fit$res, X, target = c("calib", "par", "tau"))

```

gamBiCopSelect

Selection and Maximum penalized likelihood estimation of a Generalized Additive model (gam) for the copula parameter or Kendall's tau.

Description

This function selects an appropriate bivariate copula family for given bivariate copula data using one of a range of methods. The corresponding parameter estimates are obtained by maximum penalized likelihood estimation, where each Newton-Raphson iteration is reformulated as a generalized ridge regression solved using the [mgcv](#) package.

Usage

```

gamBiCopSelect(udata, lin.covs = NULL, smooth.covs = NULL, familyset = NA,
  rotations = TRUE, familycrit = "AIC", level = 0.05, edf = 1.5,
  tau = TRUE, method = "FS", tol.rel = 0.001, n.iters = 10,
  parallel = FALSE, verbose = FALSE, select.once = TRUE, ...)

```

Arguments

udata	A matrix or data frame containing the model responses, (u1,u2) in [0,1]x[0,1]
lin.covs	A matrix or data frame containing the parametric (i.e., linear) covariates.
smooth.covs	A matrix or data frame containing the non-parametric (i.e., smooth) covariates.
familyset	(Similar to BiCopSelect from the VineCopula package) Vector of bivariate copula families to select from. If familyset = NA (default), selection among all possible families is performed. Coding of bivariate copula families: 1 Gaussian, 2 Student t, 5 Frank, 301 Double Clayton type I (standard and rotated 90 degrees), 302 Double Clayton type II (standard and rotated 270 degrees), 303 Double Clayton type III (survival and rotated 90 degrees), 304 Double Clayton type IV (survival and rotated 270 degrees), 401 Double Gumbel type I (standard and rotated 90 degrees), 402 Double Gumbel type II (standard and rotated 270 degrees), 403 Double Gumbel type III (survival and rotated 90 degrees), 404 Double Gumbel type IV (survival and rotated 270 degrees).
rotations	If TRUE, all rotations of the families in familyset are included.
familycrit	Character indicating the criterion for bivariate copula selection. Possible choices: familycrit = 'AIC' (default) or 'BIC', as in BiCopSelect from the VineCopula package.
level	Numerical; significance level of the test for removing individual predictors (default: level = 0.05).
edf	Numerical; if the estimated EDF for individual predictors is smaller than edf but the predictor is still significant, then it is set as linear (default: edf = 1.5).
tau	FALSE for a calibration function specified for the Copula parameter or TRUE (default) for a calibration function specified for Kendall's tau.
method	'FS' for Fisher-scoring (default) and 'NR' for Newton-Raphson.
tol.rel	Relative tolerance for 'FS'/'NR' algorithm.
n.iters	Maximal number of iterations for 'FS'/'NR' algorithm.
parallel	TRUE for a parallel estimation across copula families.
verbose	TRUE prints informations during the estimation.
select.once	if TRUE the GAM structure is only selected once, for the family that appears first in familyset.
...	Additional parameters to be passed to gam

Value

gamBiCopFit returns a list consisting of

res	S4 gamBiCop-class object.
method	'FS' for Fisher-scoring and 'NR' for Newton-Raphson.
tol.rel	relative tolerance for 'FS'/'NR' algorithm.
n.iters	maximal number of iterations for 'FS'/'NR' algorithm.
trace	the estimation procedure's trace.
conv	0 if the algorithm converged and 1 otherwise.

See Also

[gamBiCop](#) and [gamBiCopFit](#).

Examples

```

require(copula)
set.seed(0)

## Simulation parameters (sample size, correlation between covariates,
## Student copula with 4 degrees of freedom)
n <- 5e2
rho <- 0.9
fam <- 2
par2 <- 4

## A calibration surface depending on four variables
eta0 <- 1
calib.surf <- list(calib.lin <- function(t, Ti = 0, Tf = 1, b = 2) {
  return(-2+4*t)},
  calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
    Tm <- (Tf - Ti)/2
    a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
    return(a + b * (t - Tm)^2)},
  calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
    a <- b * (1 - 2 * Tf * pi / (f * Tf * pi +
      cos(2 * f * pi * (Tf - Ti))
      - cos(2 * f * pi * Ti)))
    return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti))/2)},
  calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
    Tm <- (Tf - Ti)/2
    a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
    return(a + b * exp(-(t - Tm)^2/(2 * s^2))))}

## 6-dimensional matrix X of covariates
covariates.distr <- mvdc(normalCopula(rho, dim = 6),
  c("unif"), list(list(min = 0, max = 1)),
  marginsIdentical = TRUE)
X <- rMvdc(n, covariates.distr)
colnames(X) <- paste("x",1:6,sep="")

## U in [0,1]x[0,1] depending on the four first columns of X
U <- condBiCopSim(fam, function(x1,x2,x3,x4) {eta0+sum(mapply(function(f,x)
  f(x), calib.surf, c(x1,x2,x3,x4)))}, X[,1:4], par2 = 4, return.par = TRUE)

## Not run:
## Selection using AIC (about 30sec on single core)
## Use parallel = TRUE to speed-up...
system.time(best <- gamBiCopSelect(U$data, smooth.covs = X))
print(best$res)
EDF(best$res) ## The first function is linear
## Plot only the smooth component
par(mfrow=c(2,2))

```

```
plot(best$res)
## End(Not run)
```

```
gamBiCopSimulate      Simulate from gamBiCop-class object
```

Description

Simulate from `gamBiCop-class` object

Usage

```
gamBiCopSimulate(object, newdata = NULL, N = NULL, return.calib = FALSE,
  return.par = FALSE, return.tau = FALSE)
```

Arguments

<code>object</code>	<code>gamBiCop-class</code> object.
<code>newdata</code>	(same as in <code>predict.gam</code> from the <code>mgcv</code> package) A matrix or data frame containing the values of the model covariates at which simulations are required. If this is not provided then simulations corresponding to the original data are returned.
<code>N</code>	sample size.
<code>return.calib</code>	should the calibration function (TRUE) be returned or not (FALSE)?
<code>return.par</code>	should the copula parameter (TRUE) be returned or not (FALSE)?
<code>return.tau</code>	should the Kendall's tau (TRUE) be returned or not (FALSE)?

Value

A list with 1 item `data`. When `N` is smaller or larger than the `newdata`'s number of rows (or the number of rows in the original data if `newdata` is not provided), then `N` observations are sampled uniformly (with replacement) among the row of `newdata` (or the rows of the original data if `newdata` is not provided).

If `return.calib = TRUE`, `return.par = TRUE` and/or `return.tau = TRUE`, then the list also contains respectively items `calib`, `par` and/or `tau`.

Examples

```
require(copula)
set.seed(1)

## Simulation parameters (sample size, correlation between covariates,
## Gaussian copula family)
n <- 5e2
rho <- 0.5
fam <- 1
```

```

## A calibration surface depending on three variables
eta0 <- 1
calib.surf <- list(
  calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
    Tm <- (Tf - Ti)/2
    a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
    return(a + b * (t - Tm)^2)},
  calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
    a <- b * (1 - 2 * Tf * pi/(f * Tf * pi +
      cos(2 * f * pi * (Tf - Ti))
      - cos(2 * f * pi * Ti)))
    return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti)/2)},
  calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
    Tm <- (Tf - Ti)/2
    a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
    return(a + b * exp(-(t - Tm)^2/(2 * s^2)))})

## 3-dimensional matrix X of covariates
covariates.distr <- mvdc(normalCopula(rho, dim = 3),
  c("unif"), list(list(min = 0, max = 1)),
  marginsIdentical = TRUE)
X <- rMvdc(n, covariates.distr)
colnames(X) <- paste("x",1:3,sep="")

## U in [0,1]x[0,1] with copula parameter depending on X
U <- condBiCopSim(fam, function(x1,x2,x3) {eta0+sum(mapply(function(f,x)
  f(x), calib.surf, c(x1,x2,x3)))}, X[,1:3], par2 = 6, return.par = TRUE)

## Merge U and X
data <- data.frame(U$data,X)
names(data) <- c(paste("u",1:2,sep=""),paste("x",1:3,sep=""))

## Model fit with penalized cubic splines (via min GCV)
basis <- c(3, 10, 10)
formula <- ~s(x1, k = basis[1], bs = "cr") +
  s(x2, k = basis[2], bs = "cr") +
  s(x3, k = basis[3], bs = "cr")
system.time(fit <- gamBiCopFit(data, formula, fam))

## Extract the gamBiCop objects and show various methods
(res <- fit$res)
EDF(res)
sim <- gamBiCopSimulate(fit$res, X)

```

Description

Constructs an object of the class `gamVine`.

Usage

```
gamVine(Matrix, model, names = NA, covariates = NA)
```

Arguments

Matrix	lower triangular $d \times d$ matrix that defines the tree structure.
model	list containing $d \times (d-1)/2$ lists with three numeric items (family, par and par2) and/or objects of the class gamBiCop .
names	vector of d names.
covariates	vector of names for the covariates.

Value

An object of the class [gamVine](#).

See Also

[gamVine](#), [RVineMatrix](#), [gamBiCop](#) [gamVineSeqFit](#), [gamVineCopSelect](#), [gamVineStructureSelect](#) and [gamVineSimulate](#).

gamVine-class

The gamVine Class

Description

`gamVine` is an S4 class to store a conditional and potentially non-simplified pair-copula construction. Objects can be created by calls of the form `new("gamVine", ...)`, or by function [gamVine](#).

Slots

Matrix	Lower triangular $d \times d$ matrix that defines the tree structure.
model	list containing $d \times (d-1)/2$ lists with three numeric items (family, par and par2) and/or gamBiCop objects.
names	vector of d names.
covariates	vector of names for the exogenous covariates.

See Also

[gamVine](#), [RVineMatrix](#), [gamBiCop](#) [gamVineSeqFit](#), [gamVineCopSelect](#), [gamVineStructureSelect](#) and [gamVineSimulate](#).

gamVineCopSelect	<i>Sequential pair-copula selection and maximum penalized likelihood estimation of a GAM-Vine model.</i>
------------------	--

Description

This function select the copula family and estimates the parameter(s) of a Generalized Additive model (GAM) Vine model, where GAMs for individual edges are specified either for the copula parameter or Kendall's tau. It solves the maximum penalized likelihood estimation for the copula families supported in this package by reformulating each Newton-Raphson iteration as a generalized ridge regression, which is solved using the [mgcv](#) package.

Usage

```
gamVineCopSelect(data, Matrix, lin.covs = NULL, smooth.covs = NULL,
  simplified = FALSE, familyset = NA, rotations = TRUE,
  familycrit = "AIC", level = 0.05, trunclevel = NA, tau = TRUE,
  method = "FS", tol.rel = 0.001, n.iters = 10, parallel = FALSE,
  verbose = FALSE, select.once = TRUE)
```

Arguments

data	A matrix or data frame containing the data in $[0,1]^d$.
Matrix	Lower triangular $d \times d$ matrix that defines the R-vine tree structure.
lin.covs	A matrix or data frame containing the parametric (i.e., linear) covariates (default: lin.covs = NULL).
smooth.covs	A matrix or data frame containing the non-parametric (i.e., smooth) covariates (default: smooth.covs = NULL).
simplified	If TRUE, then a simplified vine is fitted (which is possible only if there are exogenous covariates). If FALSE (default), then a non-simplified vine is fitted.
familyset	An integer vector of pair-copula families to select from (the independence copula MUST NOT be specified in this vector unless one wants to fit an independence vine!). The vector has to include at least one pair-copula family that allows for positive and one that allows for negative dependence. Not listed copula families might be included to better handle limit cases. If familyset = NA (default), selection among all possible families is performed. Coding of pair-copula families: 1 Gaussian, 2 Student t, 3 Clayton, 4 Gumbel, 13 Survival Clayton, 14 Survival Gumbel, 23 Rotated (90 degrees) Clayton, 24 Rotated (90 degrees) Gumbel, 33 Rotated (270 degrees) Clayton and 34 Rotated (270 degrees) Gumbel.
rotations	If TRUE, all rotations of the families in familyset are included.
familycrit	Character indicating the criterion for bivariate copula selection. Possible choices: familycrit = 'AIC' (default) or 'BIC', as in BiCopSelect from the VineCopula package.

level	Numerical; Passed to <code>gamBiCopSelect</code> , it is the significance level of the test for removing individual predictors (default: <code>level = 0.05</code>) for each conditional pair-copula.
trunclevel	Integer; level of truncation.
tau	TRUE (default) for a calibration function specified for Kendall's tau or FALSE for a calibration function specified for the Copula parameter.
method	'NR' for Newton-Raphson and 'FS' for Fisher-scoring (default).
tol.rel	Relative tolerance for 'FS'/'NR' algorithm.
n.iters	Maximal number of iterations for 'FS'/'NR' algorithm.
parallel	TRUE (default) for parallel selection of copula family at each edge or FALSE for the sequential version. for the Copula parameter.
verbose	TRUE if informations should be printed during the estimation and FALSE (default) for a silent version. from <code>mgcv</code> .
select.once	if TRUE the GAM structure is only selected once, for the family that appears first in <code>familyset</code> .

Value

`gamVineCopSelect` returns a `gamVine-class` object.

See Also

`gamVineSeqFit`, `gamVineStructureSelect`, `gamVine-class`, `gamVineSimulate` and `gamBiCopFit`.

Examples

```
require(mgcv)
set.seed(0)

## Simulation parameters
# Sample size
n <- 1e3
# Copula families
familyset <- c(1:2,301:304,401:404)
# Define a 4-dimensional R-vine tree structure matrix
d <- 4
Matrix <- c(2,3,4,1,0,3,4,1,0,0,4,1,0,0,0,1)
Matrix <- matrix(Matrix,d,d)
nnames <- paste("X", 1:d, sep = "")

## A function factory
eta0 <- 1
calib.surf <- list(
  calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
    Tm <- (Tf - Ti)/2
    a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
    return(a + b * (t - Tm)^2)},
  calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
    a <- b * (1 - 2 * Tf * pi / (f * Tf * pi +
```



```

        cos(2 * f * pi * (Tf - Ti))
        - cos(2 * f * pi * Ti)))
    return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti))/2)},
calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
  Tm <- (Tf - Ti)/2
  a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
  return(a + b * exp(-(t - Tm)^2/(2 * s^2)))})

## Create the model
# Define gam-vine model list
count <- 1
model <- vector(mode = "list", length = d*(d-1)/2)
sel <- seq(d,d^2-d, by = d)

# First tree
for (i in 1:(d-1)) {
  # Select a copula family
  family <- sample(familyset, 1)
  model[[count]]$family <- family

  # Use the canonical link and a randomly generated parameter
  if (is.element(family,c(1,2))) {
    model[[count]]$par <- tanh(rnorm(1)/2)
    if (family == 2) {
      model[[count]]$par2 <- 2+exp(rnorm(1))
    }
  } else {
    if (is.element(family,c(401:404))) {
      rr <- rnorm(1)
      model[[count]]$par <- sign(rr)*(1+abs(rr))
    } else {
      model[[count]]$par <- rnorm(1)
    }
    model[[count]]$par2 <- 0
  }
  count <- count + 1
}

# A dummy dataset
data <- data.frame(u1 = runif(1e2), u2 = runif(1e2), matrix(runif(1e2*d),1e2,d))

# Trees 2 to (d-1)
for(j in 2:(d-1)){
  for(i in 1:(d-j)){
    # Select a copula family
    family <- sample(familyset, 1)

    # Select the conditioning set and create a model formula
    cond <- nnames[sort(Matrix[(d-j+2):d,i])]
    tmpform <- paste("~",paste(paste("s(", cond, ", k=10, bs='cr')",
      sep = ""), collapse=" + "))
    l <- length(cond)
    temp <- sample(3, l, replace = TRUE)
  }
}

```

```

# Spline approximation of the true function
m <- 1e2
x <- matrix(seq(0,1,length.out=m), nrow = m, ncol = 1)
if(1 != 1){
  tmp.fct <- paste("function(x){eta0+",
                  paste(sapply(1:l, function(x)
                             paste("calib.surf[[",temp[x],"]](x[,x,""],
                                   sep="")), collapse="+"),"}",sep="")
  tmp.fct <- eval(parse(text = tmp.fct))
  x <- eval(parse(text = paste0("expand.grid(",
                                paste0(rep("x",l), collapse = ","),")",
                                collapse = "")))

  y <- apply(x,1,tmp.fct)
}else{
  tmp.fct <- function(x) eta0+calib.surf[[temp]](x)
  colnames(x) <- cond
  y <- tmp.fct(x)
}

# Estimate the gam model
form <- as.formula(paste0("y", tmpform))
dd <- data.frame(y, x)
names(dd) <- c("y", cond)
b <- gam(form, data = dd)
#plot(x[,1],(y-fitted(b))/y)

# Create a dummy gamBiCop object
tmp <- gamBiCopFit(data = data, formula = form, family = 1, n.iters = 1)$res

# Update the copula family and the model coefficients
attr(tmp, "model")$coefficients <- coefficients(b)
attr(tmp, "model")$smooth <- b$smooth
attr(tmp, "family") <- family
if (family == 2) {
  attr(tmp, "par2") <- 2+exp(rnorm(1))
}
model[[count]] <- tmp
count <- count+1
}
}

# Create the gamVineCopula object
GVC <- gamVine(Matrix=Matrix,model = model,names=nnames)
print(GVC)

## Not run:
## Simulate and fit the model
sim <- gamVineSimulate(n, GVC)
fitGVC <- gamVineSeqFit(sim, GVC, verbose = TRUE)
fitGVC2 <- gamVineCopSelect(sim, Matrix, verbose = TRUE)

## Plot the results

```

```
par(mfrow=c(3,4))
plot(GVC, ylim = c(-2.5,2.5))

plot(fitGVC, ylim = c(-2.5,2.5))

plot(fitGVC2, ylim = c(-2.5,2.5))
## End(Not run)
```

gamVineFamily*Family Matrix of an Object of the Class gamVine*

Description

Return the matrix of copula family (see [gamBiCop](#)) corresponding to the model list in the [gamVine](#) object.

Usage

```
gamVineFamily(GVC)
```

Arguments

GVC An object of the class [gamVine](#).

Value

Matrix of copula families corresponding to the model list in the [gamVine](#) object.

See Also

[gamVine](#).

gamVineNormalize*Normalize an Object of the Class gamVine*

Description

Change the R-vine matrix in the natural order, i.e. with d:1 on the diagonal

Usage

```
gamVineNormalize(GVC)
```

Arguments

GVC An object of the class [gamVine](#).

Value

The normalized [gamVine](#) object.

See Also

[gamVine](#).

gamVineSeqFit	<i>Sequential maximum penalized likelihood estimation of a GAM-Vine model.</i>
---------------	--

Description

This function estimates the parameter(s) of a Generalized Additive model (GAM) Vine model, where GAMs for individual edges are specified either for the copula parameter or Kendall's tau. It solves the maximum penalized likelihood estimation for the copula families supported in this package by reformulating each Newton-Raphson iteration as a generalized ridge regression, which is solved using the [mgcv](#) package.

Usage

```
gamVineSeqFit(data, GVC, covariates = NA, method = "FS", tol.rel = 0.001,
  n.iters = 10, verbose = FALSE)
```

Arguments

data	A matrix or data frame containing the data in $[0,1]^d$.
GVC	A gamVine object.
covariates	Vector of names for the covariates.
method	'NR' for Newton-Raphson and 'FS' for Fisher-scoring (default).
tol.rel	Relative tolerance for 'FS'/'NR' algorithm.
n.iters	Maximal number of iterations for 'FS'/'NR' algorithm.
verbose	TRUE if informations should be printed during the estimation and FALSE (default) for a silent version.
...	Additional parameters to be passed to gam from mgcv .

Value

gamVineSeqFit returns a [gamVine](#) object.

See Also

[gamVineCopSelect](#) and [gamVineStructureSelect](#)
[gamVineCopSelect](#), [gamVineStructureSelect](#), [gamVine-class](#), [gamVineSimulate](#) and [gamBiCopFit](#).

Examples

```

require(mgcv)
set.seed(0)

## Simulation parameters
# Sample size
n <- 1e3
# Copula families
familyset <- c(1:2,301:304,401:404)
# Define a 4-dimensional R-vine tree structure matrix
d <- 4
Matrix <- c(2,3,4,1,0,3,4,1,0,0,4,1,0,0,0,1)
Matrix <- matrix(Matrix,d,d)
nnames <- paste("X", 1:d, sep = "")

## A function factory
eta0 <- 1
calib.surf <- list(
  calib.quad <- function(t, Ti = 0, Tf = 1, b = 8) {
    Tm <- (Tf - Ti)/2
    a <- -(b/3) * (Tf^2 - 3 * Tf * Tm + 3 * Tm^2)
    return(a + b * (t - Tm)^2)},
  calib.sin <- function(t, Ti = 0, Tf = 1, b = 1, f = 1) {
    a <- b * (1 - 2 * Tf * pi / (f * Tf * pi +
      cos(2 * f * pi * (Tf - Ti))
      - cos(2 * f * pi * Ti)))
    return((a + b)/2 + (b - a) * sin(2 * f * pi * (t - Ti))/2)},
  calib.exp <- function(t, Ti = 0, Tf = 1, b = 2, s = Tf/8) {
    Tm <- (Tf - Ti)/2
    a <- (b * s * sqrt(2 * pi)/Tf) * (pnorm(0, Tm, s) - pnorm(Tf, Tm, s))
    return(a + b * exp(-(t - Tm)^2/(2 * s^2)))})

## Create the model
# Define gam-vine model list
count <- 1
model <- vector(mode = "list", length = d*(d-1)/2)
sel <- seq(d,d^2-d, by = d)

# First tree
for (i in 1:(d-1)) {
  # Select a copula family
  family <- sample(familyset, 1)
  model[[count]]$family <- family

  # Use the canonical link and a randomly generated parameter
  if (is.element(family,c(1,2))) {
    model[[count]]$par <- tanh(rnorm(1)/2)
    if (family == 2) {
      model[[count]]$par2 <- 2+exp(rnorm(1))
    }
  } else {
    if (is.element(family,c(401:404))) {

```

```

    rr <- rnorm(1)
    model[[count]]$par <- sign(rr)*(1+abs(rr))
  } else {
    model[[count]]$par <- rnorm(1)
  }
  model[[count]]$par2 <- 0
}
count <- count + 1
}

# A dummy dataset
data <- data.frame(u1 = runif(1e2), u2 = runif(1e2), matrix(runif(1e2*d),1e2,d))

# Trees 2 to (d-1)
for(j in 2:(d-1)){
  for(i in 1:(d-j)){
    # Select a copula family
    family <- sample(familyset, 1)

    # Select the conditioning set and create a model formula
    cond <- nnames[sort(Matrix[(d-j+2):d,i])]
    tmpform <- paste("~",paste(paste("s(", cond, ", k=10, bs='cr')",
                                   sep = "")), collapse=" + ")

    l <- length(cond)
    temp <- sample(3, l, replace = TRUE)

    # Spline approximation of the true function
    m <- 1e2
    x <- matrix(seq(0,1,length.out=m), nrow = m, ncol = 1)
    if(l != 1){
      tmp.fct <- paste("function(x){eta0+",
                      paste(sapply(1:l, function(x)
                                   paste("calib.surf[[",temp[x],"]](x[,x,")",
                                       sep="")), collapse="+"),")",sep="")
      tmp.fct <- eval(parse(text = tmp.fct))
      x <- eval(parse(text = paste0("expand.grid(",
                                   paste0(rep("x",l), collapse = ",,")"),",",
                                   collapse = ")))")

      y <- apply(x,1,tmp.fct)
    }else{
      tmp.fct <- function(x) eta0+calib.surf[[temp]](x)
      colnames(x) <- cond
      y <- tmp.fct(x)
    }

    # Estimate the gam model
    form <- as.formula(paste0("y", tmpform))
    dd <- data.frame(y, x)
    names(dd) <- c("y", cond)
    b <- gam(form, data = dd)
    #plot(x[,1],(y-fitted(b))/y)

    # Create a dummy gamBiCop object

```

```

tmp <- gamBiCopFit(data = data, formula = form, family = 1, n.iters = 1)$res

# Update the copula family and the model coefficients
attr(tmp, "model")$coefficients <- coefficients(b)
attr(tmp, "model")$smooth <- b$smooth
attr(tmp, "family") <- family
if (family == 2) {
  attr(tmp, "par2") <- 2+exp(rnorm(1))
}
model[[count]] <- tmp
count <- count+1
}
}

# Create the gamVineCopula object
GVC <- gamVine(Matrix=Matrix,model = model,names=nnames)
print(GVC)

## Not run:
## Simulate and fit the model
sim <- gamVineSimulate(n, GVC)
fitGVC <- gamVineSeqFit(sim, GVC, verbose = TRUE)
fitGVC2 <- gamVineCopSelect(sim, Matrix, verbose = TRUE)

## Plot the results
dev.off()
par(mfrow=c(3,4))
plot(GVC, ylim = c(-2.5,2.5))

plot(fitGVC, ylim = c(-2.5,2.5))

plot(fitGVC2, ylim = c(-2.5,2.5))
## End(Not run)

```

gamVineSimulate

Simulation from a [gamVine-class](#) object

Description

Simulation from a [gamVine-class](#) object

Usage

```
gamVineSimulate(n, GVC, U = NULL, newdata = NULL)
```

Arguments

n number of d-dimensional observations to simulate.
GVC A [gamVine](#) object.

U	If not NULL, U is an (N,d)-matrix of U[0,1] random variates to be transformed to the copula sample.
newdata	If not NULL, which is mandatory when the attribute covariates from GVC is not NA, newdata is a data frame containing the values of the model covariates at which simulations are required.

Value

A matrix of data simulated from the given `gamVine` object.

Examples

```
require(VineCopula)

## Example adapted from RVineSim

## Define 5-dimensional R-vine tree structure matrix
Matrix <- c(5, 2, 3, 1, 4,
            0, 2, 3, 4, 1,
            0, 0, 3, 4, 1,
            0, 0, 0, 4, 1,
            0, 0, 0, 0, 1)
Matrix <- matrix(Matrix, 5, 5)

## Define R-vine pair-copula family matrix
family <- c(0, 1, 3, 4, 4,
            0, 0, 3, 4, 1,
            0, 0, 0, 4, 1,
            0, 0, 0, 0, 3,
            0, 0, 0, 0, 0)
family <- matrix(family, 5, 5)

## Define R-vine pair-copula parameter matrix
par <- c(0, 0.2, 0.9, 1.5, 3.9,
        0, 0, 1.1, 1.6, 0.9,
        0, 0, 0, 1.9, 0.5,
        0, 0, 0, 0, 4.8,
        0, 0, 0, 0, 0)
par <- matrix(par, 5, 5)

## Define second R-vine pair-copula parameter matrix
par2 <- matrix(0, 5, 5)

## Define RVineMatrix object
RVM <- RVineMatrix(Matrix = Matrix, family = family,
                  par = par, par2 = par2,
                  names = c("V1", "V2", "V3", "V4", "V5"))

## Convert to gamVine object
GVC <- RVM2GVC(RVM)

## U[0,1] random variates to be transformed to the copula sample
```



```

n <- 1e2
d <- 5
U <- matrix(runif(n*d), nrow = n)

## The output of gamVineSimulate correspond to that of RVineSim
sampleRVM <- RVineSim(n, RVM,U)
sampleGVC <- gamVineSimulate(n, GVC,U)
all.equal(sampleRVM, sampleGVC)

## Fit the two models and compare the estimated parameter
fitRVM <- RVM2GVC(RVineSeqEst(sampleRVM,RVM))
fitGVC <- gamVineSeqFit(sampleGVC,GVC)
all.equal(simplify2array(attr(fitRVM, "model")),
simplify2array(attr(fitGVC, "model")))

```

gamVineStructureSelect

Structure selection and estimation of a GAM-Vine model.

Description

This function select the structure and estimates the parameter(s) of a Generalized Additive model (GAM) Vine model, where GAMs for individual edges are specified either for the copula parameter or Kendall's tau. It solves the maximum penalized likelihood estimation for the copula families supported in this package by reformulating each Newton-Raphson iteration as a generalized ridge regression, which is solved using the [mgcv](#) package.

Usage

```

gamVineStructureSelect(udata, lin.covs = NULL, smooth.covs = NULL,
  simplified = TRUE, type = 0, familyset = NA, rotations = TRUE,
  familycrit = "AIC", treecrit = "tau", level = 0.05, trunclevel = NA,
  tau = TRUE, method = "FS", tol.rel = 0.001, n.iters = 10,
  parallel = FALSE, verbose = FALSE, select.once = TRUE)

```

Arguments

udata	A matrix or data frame containing the data in $[0,1]^d$.
lin.covs	A matrix or data frame containing the parametric (i.e., linear) covariates (default: lin.covs = NULL).
smooth.covs	A matrix or data frame containing the non-parametric (i.e., smooth) covariates (default: smooth.covs = NULL).
simplified	If TRUE (default), then a simplified vine is fitted (which is possible only if there are exogenous covariates). If FALSE, then a non-simplified vine is fitted.
type	type = 0 (default) for a R-Vine and type = 1 for a C-Vine.

familyset	An integer vector of pair-copula families to select from (the independence copula MUST NOT be specified in this vector unless one wants to fit an independence vine!). Not listed copula families might be included to better handle limit cases. If familyset = NA (default), selection among all possible families is performed. Coding of pair-copula families: 1 Gaussian, 2 Student t, 5 Frank, 301 Double Clayton type I (standard and rotated 90 degrees), 302 Double Clayton type II (standard and rotated 270 degrees), 303 Double Clayton type III (survival and rotated 90 degrees), 304 Double Clayton type IV (survival and rotated 270 degrees), 401 Double Gumbel type I (standard and rotated 90 degrees), 402 Double Gumbel type II (standard and rotated 270 degrees), 403 Double Gumbel type III (survival and rotated 90 degrees), 404 Double Gumbel type IV (survival and rotated 270 degrees).
rotations	If TRUE, all rotations of the families in familyset are included.
familycrit	Character indicating the criterion for bivariate copula selection. Possible choices: familycrit = 'AIC' (default) or 'BIC', as in BiCopSelect from the VineCopula package.
treecrit	Character indicating how pairs are selected in each tree. treecrit = "tau" uses the maximum spanning tree of the Kendall's tau (i.e., the tree of maximal overall dependence), treecrit = "rho" uses the Spearman's rho.
level	Numerical; Passed to gamBiCopSelect , it is the significance level of the test for removing individual predictors (default: level = 0.05) for each conditional pair-copula.
trunclevel	Integer; level of truncation.
tau	TRUE (default) for a calibration function specified for Kendall's tau or FALSE for a calibration function specified for the Copula parameter.
method	'NR' for Newton-Raphson and 'FS' for Fisher-scoring (default).
tol.rel	Relative tolerance for 'FS'/'NR' algorithm.
n.iters	Maximal number of iterations for 'FS'/'NR' algorithm.
parallel	TRUE (default) for parallel selection of copula family at each edge or FALSE for the sequential version. for the Copula parameter.
verbose	TRUE if informations should be printed during the estimation and FALSE (default) for a silent version. from mgcv .
select.once	if TRUE the GAM structure is only selected once, for the family that appears first in familyset.

Value

gamVineSeqFit returns a [gamVine-class](#) object.

See Also

[gamVineSeqFit](#), [gamVineCopSelect](#), [gamVine-class](#), [gamVineSimulate](#) and [gamBiCopSelect](#).

Examples

```

require(VineCopula)
set.seed(0)

## An example with a 3-dimensional GAM-Vine

# Sample size
n <- 1e3

# Define a R-vine tree structure matrix
d <- 3
Matrix <- c(2,3,1,0,3,1,0,0,1)
Matrix <- matrix(Matrix,d,d)
nnames <- paste("x", 1:d, sep = "")

# Copula families for each edge
fam <- c(301,401,1)

# Parameters for the first tree (two unconditional copulas)
par <- c(1,2)

# Pre-allocate the GAM-Vine model list
count <- 1
model <- vector(mode = "list", length = d*(d-1)/2)

# The first tree contains only the two unconditional copulas
for (i in 1:(d-1)) {
  model[[count]] <- list(family = fam[count], par = par[count], par2 = 0)
  count <- count + 1
}

# The second tree contains a unique conditional copula
# In this first example, we take a linear calibration function (10*x-5)

# Set-up a dummy dataset
tmp <- data.frame(u1 = runif(1e2), u2 = runif(1e2), x1 = runif(1e2))

# Set-up an arbitrary linear model for the calibration function
model[[count]] <- gamBiCopFit(tmp, ~ x1, fam[count])$res

# Update the coefficients of the model
attr(model[[count]],"model")$coefficients <- c(-5, 10)

# Define gamVine object
GVC <- gamVine(Matrix = Matrix, model = model, names = nnames)
GVC

## Not run:
# Simulate new data
simData <- data.frame(gamVineSimulate(n, GVC))
colnames(simData) <- nnames

```

```
# Fit data using sequential estimation assuming true model known
summary(fitGVC <- gamVineSeqFit(simData, GVC))

# Fit data using structure selection and sequential estimation
summary(fitGVC2 <- gamVineStructureSelect(simData, simplified = FALSE))
## End(Not run)
```

logLik.gamBiCop	<i>Extract the Log-likelihood from a gamBiCop Object</i>
-----------------	--

Description

Function to extract the log-likelihood from an object of the class `gamBiCop` (note that the models are usually fitted by penalized likelihood maximization). This function is used by [AIC](#) and [BIC](#).

Usage

```
## S4 method for signature 'gamBiCop'
logLik(object, ...)
```

Arguments

object	An object of the class <code>gamBiCop</code> .
...	un-used in this class

Value

Standard logLik object: see [logLik](#).

See Also

[AIC](#) and [BIC](#).

nobs.gamBiCop	<i>Extract the Number of Observations from gamBiCop Object</i>
---------------	--

Description

Extract the number of 'observations' from a model fit. This is principally intended to be used in computing the BIC (see [AIC](#)).

Usage

```
## S4 method for signature 'gamBiCop'
nobs(object, ...)
```

Arguments

object An object of the class [gamBiCop](#).
 ... un-used in this class

Value

A single number, normally an integer.

See Also

[AIC](#) and [BIC](#).

plot.gamBiCop	<i>Plot a gamBiCop Object</i>
---------------	-------------------------------

Description

Plot from an object of the class [gamBiCop](#). The function is based on (see [plot.gam](#) from [mgcv](#)).

Usage

```
## S4 method for signature 'gamBiCop,ANY'
plot(x, y, ...)
```

Arguments

x An object of the class [gamBiCop](#).
 y Not used with this class.
 ... additional arguments to be passed to [plot.gam](#).

Value

This function simply generates plots.

See Also

[plot.gam](#) from [mgcv](#)).

plot.gamVine	<i>Plot an Object of the Class gamVine</i>
--------------	--

Description

Plot an object of the class [gamVine](#). The function is based on (see [plot.gam](#) from [mgcv](#)).

Usage

```
## S4 method for signature 'gamVine,ANY'
plot(x, y, ...)
```

Arguments

x	An object of the class gamVine .
y	Not used with this class.
...	additional arguments to be passed to plot.gam .

Value

This function simply generates plots.

See Also

[plot.gam](#) from [mgcv](#)).

RVM2GVC	<i>Transform an Object of the Class R-Vine into an Object of the Class gamVine</i>
---------	--

Description

Transform an object of the class [RVineMatrix](#) into an object of the class [gamVine](#).

Usage

```
RVM2GVC(RVM)
```

Arguments

RVM	An object of the class RVineMatrix .
-----	--

Value

An object of the class [gamVine](#).

See Also

[RVineMatrix](#) and [gamVine](#).

summary.gamBiCop	<i>Summary for a gamBiCop Object</i>
------------------	--------------------------------------

Description

Takes a [gamBiCop](#) object and produces various useful summaries from it.

Usage

```
## S4 method for signature 'gamBiCop'
summary(object)
```

Arguments

object	An object of the class gamBiCop .
...	unused in this class

Value

A useful summary (see [summary.gam](#) from [mgcv](#) for more details).

See Also

[summary.gam](#) from [mgcv](#)

summary.gamVine	<i>Summary for an Object of the Class gamVine</i>
-----------------	---

Description

Takes an object of the class [gamVine](#) and produces various useful summaries from it.

Usage

```
## S4 method for signature 'gamVine'
summary(object)
```

Arguments

object	An object of the class gamVine .
...	unused in this class

Value

A useful summary (see [summary.gam](#) from [mgcv](#) for more details).

See Also

[summary.gam](#) from [mgcv](#)

Index

*Topic **package**

- gamCopula-package, 2
- AIC, 9, 44, 45
- AIC, gamBiCop-method (AIC.gamBiCop), 8
- AIC.gamBiCop, 8
- BIC, 9, 44, 45
- BIC, gamBiCop-method (BIC.gamBiCop), 9
- BIC.gamBiCop, 9
- BiCopEta2Par, 9, 10, 11
- BiCopPar2Eta, 10
- BiCopPar2Tau, 10, 11
- BiCopSelect, 26, 31, 42
- BiCopTau2Par, 10, 11
- condBiCopSim, 11
- dim, gamVine-method (dim.gamVine), 13
- dim.gamVine, 13
- EDF, 14
- formula, gamBiCop-method
(formula.gamBiCop), 15
- formula.gam, 15, 18
- formula.gamBiCop, 15
- gam, 15, 16, 18, 19, 26, 36
- gam.models, 18
- gamBiCop, 8, 9, 14, 15, 15, 16, 17, 19, 22, 24,
27, 30, 35, 44, 45, 47
- gamBiCop-class, 16, 28
- gamBiCopCDF, 16
- gamBiCopFit, 12, 16, 18, 24, 27, 32, 36
- gamBiCopPDF, 21
- gamBiCopPredict, 16, 17, 22, 23
- gamBiCopSelect, 25, 32, 42
- gamBiCopSimulate, 12, 16, 19, 28
- gamCopula (gamCopula-package), 2
- gamCopula-package, 2
- gamCopula.package (gamCopula-package), 2
- gamObject, 14–16
- gamVine, 14, 29, 29, 30, 35, 36, 39, 40, 46, 47
- gamVine-class, 30, 39
- gamVineCopSelect, 30, 31, 36, 42
- gamVineFamily, 35
- gamVineNormalize, 35
- gamVineSeqFit, 30, 32, 36, 42
- gamVineSimulate, 30, 32, 36, 39, 42
- gamVineStructureSelect, 30, 32, 36, 41
- logLik, 44
- logLik, gamBiCop-method
(logLik.gamBiCop), 44
- logLik.gamBiCop, 44
- mgcv, 3, 15–19, 22–25, 28, 31, 32, 36, 41, 42,
45–48
- nobs, gamBiCop-method (nobs.gamBiCop), 44
- nobs.gamBiCop, 44
- plot, gamBiCop, ANY-method
(plot.gamBiCop), 45
- plot, gamVine, ANY-method (plot.gamVine),
46
- plot.gam, 45, 46
- plot.gamBiCop, 45
- plot.gamVine, 46
- predict.gam, 17, 22–24, 28
- RVineMatrix, 30, 46, 47
- RVM2GVC, 46
- summary, gamBiCop-method
(summary.gamBiCop), 47
- summary, gamVine-method
(summary.gamVine), 47
- summary.gam, 47, 48
- summary.gamBiCop, 47
- summary.gamVine, 47

VineCopula, [3](#), [10](#), [11](#), [26](#), [31](#), [42](#)