

Package ‘copBasic’

August 31, 2026

Title General Bivariate Copula Theory and Many Utility Functions

Version 2.2.16

Date 2026-08-30

Description Extensive functions for bivariate copula (bicopula) computations and related operations for bicopula theory. The lower, upper, product, and select other bicopula are implemented along with operations including the diagonal, survival copula, dual of a copula, co-copula, and numerical bicopula density. Level sets, horizontal and vertical sections are supported. Numerical derivatives and inverses of a bicopula are provided through which simulation is implemented. Bicopula composition, convex combination, asymmetry extension, and products also are provided. Support extends to the Kendall Function as well as the Lmoments thereof. Kendall Tau, Spearman Rho and Footrule, Gini Gamma, Blomqvist Beta, Hoeffding Phi, Schweizer-Wolff Sigma, tail dependency, tail order, skewness, and bivariate Lmoments are implemented, and positive/negative quadrant dependency, left (right) increasing (decreasing) are available. Other features include Kullback-Leibler Divergence, Vuong Procedure, spectral measure, and Lcomoments for fit and inference, Lcomoment ratio diagrams, maximum likelihood, and AIC, BIC, and RMSE for goodness-of-fit.

Maintainer William Asquith <william.asquith@ttu.edu>

Repository CRAN

Depends R (>= 4.0.0)

Imports lmomco, mvtnorm, randtoolbox

Suggests copula

License GPL-2

NeedsCompilation no

Author William Asquith [aut, cre] (ORCID:
<<https://orcid.org/0000-0002-7400-1861>>)

Date/Publication 2026-08-31 15:40:10 UTC

Contents

copBasic-package	5
aicCOP	15
AMHcop	17

asCOP	20
bicCOP	22
bicoploc	24
bilmoms	30
blomatrixCOP	36
blomCOP	40
blomCOPss	43
breveCOP	46
CIRCcop	49
CLcop	52
coCOP	53
composite1COP	55
composite2COP	57
composite3COP	60
convex2COP	62
convexCOP	64
COP	65
copBasic.fitpara	70
COPinv	73
COPinv2	74
densityCOP	76
densityCOPplot	78
derCOP	81
derCOP2	83
derCOPinv	85
derCOPinv2	89
diagCOP	90
diagCOPatf	92
duCOP	94
EMPIRcop	97
EMPIRcopdf	102
EMPIRgrid	103
EMPIRgridder	107
EMPIRgridder2	109
EMPIRgridderinv	110
EMPIRgridderinv2	112
EMPIRgrid_fast	113
EMPIRgrid_lkup	115
EMPIRmed.regress	116
EMPIRmed.regress2	117
EMPIRqua.regress	118
EMPIRqua.regress2	124
EMPIRsim	125
EMPIRsimv	127
EuvCOP	129
EvuCOP	131
FGMcop	134
footCOP	137

FRcop	140
FRECHETcop	143
gEVcop	145
GHcop	147
giniCOP	153
GLcop	157
glueCOP	160
gridCOP	162
HAIRPINcop	164
hoefCOP	165
HRcop	169
isCOP.LTD	171
isCOP.permsym	172
isCOP.PQD	173
isCOP.radsym	176
isCOP.RTI	179
isfuncCOP	181
JOcopB5	182
joeskewCOP	184
joint.curvesCOP	189
joint.curvesCOP2	193
jointCOP	195
JOMAcop	200
kfuncCOP	203
kfuncCOPinv	211
kfuncCOPlmoms	213
kfuncCOP_Pd	216
kullCOP	218
lcomCOP	222
lcomCOPpv	226
LCOMDIA_GH2cop	229
LCOMDIA_GH3cop	232
LCOMDIA_ManyCops	233
lcomoms2.ABcop2parameter	238
lcomoms2.ABKGcop2parameter	242
level.curvesCOP	245
level.curvesCOP2	247
level.setCOP	248
level.setCOP2	249
LMRuvCOP	250
LMRvuCOP	251
LzCOPpermsym	254
M	260
med.regressCOP	261
med.regressCOP2	264
mleCOP	265
MOcop	270
M_N5p12b	272

N4212cop	274
N4220cop	275
NORMcop	276
ORDSUMcop	278
ORDSUWcop	284
P	286
PARETOcop	287
PLACKETTcop	289
PLACKETTpar	291
PLACKETTsim	293
prod2COP	294
psepolar	297
PSP	299
qua.regressCOP	300
qua.regressCOP.draw	302
qua.regressCOP2	304
RAYcop	305
ReineckeWell266	307
ReineckeWells	308
RFcop	309
rhobevCOP	311
rhoCOP	313
rmseCOP	316
rN4220cop	317
sectionCOP	318
semicorCOP	320
simcomposite3COP	323
simcompositeCOP	325
simCOP	327
simCOPmicro	329
spectralmeas	333
stabt看depf	337
statTn	340
surCOP	343
surfuncCOP	345
tailconCOP	347
taildepCOP	348
tailordCOP	353
tauCOP	355
Tcop	358
tEVcop	361
TRIcop	363
uvmoms	364
vuongCOP	366
W	372
wolfCOP	374
wolfCOPsamc	379
wolfCOPtest	384

wolfCOPtest_check	391
wolfCOPtest_data_smlsam	397
W_N5p12a	399
Index	401

copBasic-package	<i>Basic Theoretical Copula, Empirical Copula, and Various Utility Functions</i>
------------------	--

Description

The **copBasic** package is oriented around *bivariate copula theory* and mathematical operations closely follow the recommended texts of Nelsen (2006) and Joe (2014) as well as select other references. Another recommended text is Salvadori *et al.* (2007) and Hofert *et al.* (2018) and are cited herein, but about half of that excellent book concerns univariate applications. The primal objective of **copBasic** is to provide a study of numerous results shown by authoritative texts on copulas. It is intended that the package will help other copula students in self study, potential course work, and applied circumstances.

Notes on copulas that are supported. The author has focused on pedagogical aspects of copulas, and this package is a *diary* of sorts that began in fall 2008. Originally, the author did not implement many copulas in the **copBasic** in order to deliberately avoid redundancy to that support such as it exists on the R CRAN. Though as time has progressed, other copulas have been added based on needs of a user community, needs to show some concepts in the general theory, or test algorithms. For example, the Clayton copula ([CLcop](#)) was a late arriving addition the package (*c.*2017), which was added to assist a specific user; the Frank copula ([FRcop](#)) was added in December 2024 because of an inquiry on using **copBasic** for *vine copula* (see *vine example* under [FRcop](#)). The Normal ([NORMcop](#)), Marshall–Olkin ([MOcop](#)), and t-Student copulas ([Tcop](#)) were added in summer 2025 because of an inquiry on meanings and interpretive properties of L-comoments.

The language and vocabulary of copulas are formidable even within the realm of *bivariate* or *bi-copula* as design basis for the package. Often “vocabulary” words are often emphasized in *italics*, which is used extensively and usually near the opening of function-by-function documentation to identify vocabulary words, such as *survival copula* (see [surCOP](#)). This syntax tries to mimic and accentuate the terminology in Nelsen (2006), Joe (2014), and generally most other texts. Italics are used to draw connections between concepts.

In conjunction with the summary of functions in **copBasic-package**, the extensive cross referencing to functions and expansive keyword indexing should be beneficial. The author had no experience with copulas prior to a chance happening upon Nelsen (2006) in *c.*2008. The **copBasic** package is a personal *tour de force* in self-guided learning. Over time, this package and user’s manual have been helpful to many others.

Helpful Navigation of Copulas Implemented in the copBasic Package

Some entry points to the copulas implemented are listed in the **Table of Copulas**:

Name	Symbol	Function	Concept
Lower-bounds copula	$\mathbf{W}(u, v)$	W	copula
Independence copula	$\mathbf{\Pi}(u, v)$	P	copula
Upper-bounds copula	$\mathbf{M}(u, v)$	M	copula
Fréchet Family copula	$\mathbf{FF}(u, v)$	FRECHETcop	copula
Ali–Mikhail–Haq copula	$\mathbf{AMH}(u, v)$	AMHcop	copula
Clayton copula	$\mathbf{CL}(u, v)$	CLcop	copula
Copula of uniform circle	$\mathbf{CIRC}(u, v)$	CIRCcop	copula
Farlie–Gumbel–Morgenstern (generalized)	$\mathbf{FGM}(u, v)$	FGMcop	copula
Frank	$\mathbf{FR}(u, v)$	FRcop	copula
Galambos copula	$\mathbf{GL}(u, v)$	GLcop	copula
Gumbel–Hougaard copula	$\mathbf{GH}(u, v)$	GHcop	copula
Hairpin copula	$\mathbf{HAIRPIN}(u, v)$	HAIRPINcop	copula
Hüsler–Reiss copula	$\mathbf{HR}(u, v)$	HRcop	copula
Joe B5 (the “Joe”) copula	$\mathbf{B5}(u, v)$	J0copB5	copula
Joe–Ma copula	$\mathbf{JOMA}(u, v)$	JOMAcop	copula
Marshall–Olkin copula	$\mathbf{MO}(u, v)$	M0cop	copula
Nelsen eq.4-2-12 copula	$\mathbf{N4212cop}(u, v)$	N4212cop	copula
Nelsen eq.4-2-20 copula	$\mathbf{N4220cop}(u, v)$	N4220cop	copula
Nelsen eq.4-2-20 copula (reflected)	$\mathbf{rN4220cop}(u, v)$	rN4220cop	copula
Normal copula (refer also to Tcop)	$\mathbf{NORM}(u, v)$	NORMcop	copula
Ordinal Sums by Copula	$\mathbf{C}_{\mathcal{J}}(u, v)$	ORDSUMcop	copula
Ordinal Sums by W-Copula	$\mathbf{C}_{\mathcal{J}}(u, v)$	ORDSUWcop	copula
Pareto copula	$\mathbf{PA}(u, v)$	PAcop	copula
Plackett copula	$\mathbf{PL}(u, v)$	PLcop	copula
PSP copula	$\mathbf{PSP}(u, v)$	PSP	copula
Raftery copula	$\mathbf{RF}(u, v)$	RFcop	copula
Rayleigh copula	$\mathbf{RAY}(u, v)$	RAYcop	copula
g-EV copula (Gaussian extreme value)	$\mathbf{gEV}(u, v)$	gEVcop	copula
t-EV copula (t-distribution extreme value)	$\mathbf{tEV}(u, v)$	tEVcop	copula
t-Student copula (refer also to NORMcop)	$\mathbf{T}(u, v)$	Tcop	copula
Triangular copula	$\mathbf{TRI}(u, v)$	TRICop	copula

A few comments on notation herein are needed. A bold math typeface is used to represent a copula function or family such as $\mathbf{\Pi}$ (see [P](#)) for the *independence copula*. The syntax $\mathcal{R} \times \mathcal{R} \equiv \mathcal{R}^2$ denotes the orthogonal domain of two real numbers, and $[0, 1] \times [0, 1] \equiv \mathcal{I} \times \mathcal{I} \equiv \mathcal{I}^2$ denotes the orthogonal domain on the unit square of probabilities. Limits of integration $[0, 1]$ or $[0, 1]^2$ involving copulas are thus shown as $\mathcal{I}$ and $\mathcal{I}^2$, respectively.

Random variables X and Y respectively denote the horizontal and vertical directions in $\mathcal{R}^2$. Their probabilistic counterparts are uniformly distributed random variables on $[0, 1]$, are respectively denoted as U and V , and necessarily also are the respective directions in $\mathcal{I}^2$ (U denotes the horizontal, V denotes the vertical). Often realizations of these random variables are respectively x and y for X and Y and u and v for U and V . There is an obvious difference between nonexceedance probability F and its complement, which is exceedance probability defined as $1 - F$. Both u and v herein are in nonexceedance probability. Arguments to many functions herein are $u = u$ and $v = v$ and are almost *exclusively nonexceedance* but there are instances for which the probability arguments are u

$$= 1 - u = u' \text{ and } v = 1 - v = v'.$$

Bivariate Association — Several of the functions listed above are measures of *bivariate association*. Two of the measures (*Kendall Tau*, [tauCOP](#); *Spearman Rho*, [rhoCOP](#)) are widely known. R provides native support for their sample estimation of course (`stats::cor()`), but each function can be used to call the `cor()` function for parallelism to the other measures of this package. The other measures (*Blomqvist Beta*, *Gini Gamma*, *Hoeffding Phi*, *Schweizer–Wolff Sigma*, *Spearman Footrule*) support sample estimation by specially formed calls to their respective functions: [blomCOP](#), [giniCOP](#), [hoefCOP](#), [wolfCOP](#), and [footCOP](#). *Gini Gamma* ([giniCOP](#)) documentation (also [joeskewCOP](#)) shows extensive use of theoretical and sample computations for these and other functions.

Helpful Navigation of the copBasic Package

Some other entry points into the package are listed in the following table:

Name	Symbol	Function	Concept
Copula	$\mathbf{C}(u, v)$	COP	copula theory
...	$\mathbf{C}(u, v)$	COP(..., reflect=*)	reflection
...	$\mathbf{C}(u, v)$	COP(..., reflect=*)	rotation
Survival copula	$\hat{\mathbf{C}}(u', v')$	surCOP	copula theory
Joint survival function	$\overline{\mathbf{C}}(u, v)$	surfuncCOP	copula theory
Co-copula	$\mathbf{C}^*(u', v')$	coCOP	copula theory
Dual of a copula	$\tilde{\mathbf{C}}(u, v)$	duCOP	copula theory
Primary copula diagonal	$\delta(t)$	diagCOP	copula theory
Secondary copula diagonal	$\delta^*(t)$	diagCOP	copula theory
Inverse copula diagonal	$\delta^{(-1)}(f)$	diagCOPatf	copula theory
Joint probability	—	jointCOP	copula theory
Bivariate L-moments	$\delta_{k;\mathbf{C}}^{[\dots]}$	bilmoms and lcomCOP	bivariate moments
Bivariate L-comoments	$\tau_{k;\mathbf{C}}^{[\dots]}$	bilmoms and lcomCOP	bivariate moments
Blomqvist Beta	$\beta_{\mathbf{C}}$	blomCOP	bivariate association
Gini Gamma	$\gamma_{\mathbf{C}}$	giniCOP	bivariate association
Hoeffding Phi	$\Phi_{\mathbf{C}}$	hoefCOP	bivariate association
Nu-Skew	$\nu_{\mathbf{C}}$	nuskewCOP	bivariate moments
Nu-Star (skew)	$\nu_{\mathbf{C}}^*$	nustarCOP	bivariate moments
Lp distance to independence	$\Phi_{\mathbf{C}} \rightarrow L_p$	LpCOP	bivariate association
Permutation-Mu	$\mu_{\infty\mathbf{C}}^{\text{permsym}}$	LzCOPpermsym	permutation asymmetry
Kendall Tau	$\tau_{\mathbf{C}}$	tauCOP	bivariate association
Kendall Measure	$K_{\mathbf{C}}(z)$	kmeasCOP	copula theory
Kendall Function	$F_K(z)$	kfuncCOP	copula theory
Kendall Function ($d\text{-}\mathbf{P}$)	$F_K(z; \mathbf{P}_d)$	kfuncCOP_Pd	copula theory
Inverse Kendall Function	$F_K^{(-1)}(z)$	kfuncCOPinv	copula theory
An L-moment of $F_K(z)$	$\lambda_r(F_K)$	kfuncCOPlmom	L-moment theory
L-moments of $F_K(z)$	$\lambda_r(F_K)$	kfuncCOPlmoms	L-moment theory
Semi-correlations (negatives)	$\rho_N^-(a)$	semicorCOP	bivariate tail association
Semi-correlations (positives)	$\rho_N^+(a)$	semicorCOP	bivariate tail association
Spearman Footrule	$\psi_{\mathbf{C}}$	footCOP	bivariate association
Spearman Rho	$\rho_{\mathbf{C}}$	rhoCOP	bivariate association

Schweizer–Wolff Sigma	σ_C	wolfCOP	bivariate association
...	$\hat{\sigma}_C$	wolfCOPsamc	bivariate association
...	$\sigma_C = \Pi$	wolfCOPtest	bivariate association
Density of a copula	$c(u, v)$	densityCOP	copula density
Density visualization	--	densityCOPplot	copula density
Parametric copulatic surface	--	gridCOP	copulatic surface
Parametric simulation	--	simCOP or rCOP	copula simulation
Parametric simulation	--	simCOPmicro	copula simulation
Maximum likelihood	$\mathcal{L}(\Theta_d)$	mleCOP	copula fitting
Akaike information criterion	AIC_C	aicCOP	goodness-of-fit
Bayesian information crit.	BIC_C	bicCOP	goodness-of-fit
Root mean square error	$RMSE_C$	rmseCOP	goodness-of-fit
Another goodness-of-fit	T_n	statTn	goodness-of-fit

Some entry points into the package for *empirical copula* are listed in the following table:

Name	Symbol	Function	Concept
Empirical copula	$C_n(u, v)$	EMPIRcop	copula
Empirical simulation	--	EMPIRsim	copula simulation
Empirical simulation	--	EMPIRsimv	copula simulation
Empirical (legacy) surface	$\ddot{C}_n(u, v)$	EMPIRgrid	copulatic surface
Empirical (fast) surface	$\ddot{C}_n(u, v)$	EMPIRgrid_fast	copulatic surface
Empirical copula (grid lookup)	$\ddot{C}_n(u, v)$	EMPIRgrid_lkup	copulatic surface

Goodness-of-fit — Concerning *goodness-of-fit* and although not quite the same as copula properties (such as “correlation”) per se as the coefficients aforementioned in the prior paragraph, three goodness-of-fit metrics of a copula compared to the empirical copula, which are all based the *mean square error* (MSE), are [aicCOP](#), [bicCOP](#), and [rmseCOP](#). This triad of functions is useful for making decisions on whether a copula is more favorable than another to a given dataset. However, because they are genetically related by using MSE and if these are used for copula fitting by minimization, the fits will be identical. A statement of “not quite the same” is made because the previously described copula properties are generally defined as types of deviations from other copulas (such as [P](#)). Another goodness-of-fit statistic is [statTn](#), which is based on magnitude summation of fitted copula difference from the empirical copula. These four ([aicCOP](#), [bicCOP](#), [rmseCOP](#), and [statTn](#)) collectively are relative simple and readily understood measures. These bulk sample statistics are useful, but generally thought to not capture the nuances of tail behavior ([semicorCOP](#) and [taildepCOP](#) might be useful).

Bivariate Skewness and L-comoments — *Bivariate skewness* measures are supported in the functions [joeskewCOP](#) ([nuskewCOP](#) and [nustarCOP](#)) and [uvlmoms](#) ([uvskew](#)). Extensive discussion and example computations of bivariate skewness are provided in the [joeskewCOP](#) documentation. Lastly, so-called *bivariate L-moments* and *bivariate L-comoments* of a copula are directly computable in [bilmoms](#) (though that function using Monte Carlo integration is deprecated) and [lcomCOP](#) (direct numerical integration). Function [lcomCOP](#) is the theoretical counterpart to the *sample L-comoments* provided in the **lmomco** package by `lmomco::lcomoms2()` function. Parameter estimation by

L-comoments is demonstrated for the **MOcop** under its **Examples** and for a potentially complex **composite1COP** within **Examples** of **lcomCOP**. Demonstrations of *L-comoment ratio diagrams* and construction methods for various copula is seen in **Examples** within **LCOMDIA_ManyCops** (a collection of vary many 1-parameter copula driven by Spearman Rho, **rhoCOP**), **LCOMDIA_GH2cop** (2-parameter Gumbel–Hougaard copula, **GHcop**), **LCOMDIA_GH3cop** (3-parameter Gumbel–Hougaard copula, **GHcop**), and **ORDSUMcop** (ordinal sums of M-copula).

Conditional Simulation — Random simulation methods by several functions are listed in the previous table (**simCOPmicro**, **simCOP**). The **copBasic** package explicitly uses only *conditional simulation* also known as the *conditional distribution method* for random variate generation following Nelsen (2006, pp. 40–41). The *numerical derivatives* (**derCOP** and **derCOP2**) and their *inversions* (**derCOPinv** and **derCOPinv2**) (next table) represent the foundation of the conditional simulation. There are other methods in the literature and available in other R packages, and a comparison of some methods is made in the **Examples** section of the Gumbel–Hougaard copula (**GHcop**).

Several functions in **copBasic** make the distinction between V with respect to (*wrt*) U and U wrt V , and a guide for the nomenclature involving *wrt* distinctions is listed in the following table:

Name	Symbol	Function	Concept
Copula inversion	$V \text{ wrt } U$	COPinv	copula operator
Copula inversion	$U \text{ wrt } V$	COPinv2	copula operator
Copula derivative	$\delta \mathbf{C} / \delta u$	derCOP	copula operator
Copula derivative	$\delta \mathbf{C} / \delta v$	derCOP2	copula operator
Copula derivative inversion	$V \text{ wrt } U$	derCOPinv	copula operator
Copula derivative inversion	$U \text{ wrt } V$	derCOPinv2	copula operator
Joint curves	$t \mapsto \mathbf{C}(u = U, v)$	joint.curvesCOP	copula theory
Joint curves	$t \mapsto \mathbf{C}(u, v = V)$	joint.curvesCOP2	copula theory
Level curves	$t \mapsto \mathbf{C}(u = U, v)$	level.curvesCOP	copula theory
Level curves	$t \mapsto \mathbf{C}(u, v = V)$	level.curvesCOP2	copula theory
Level set	$V \text{ wrt } U$	level.setCOP	copula theory
Level set	$U \text{ wrt } V$	level.setCOP2	copula theory
Median regression	$V \text{ wrt } U$	med.regressCOP	copula theory
Median regression	$U \text{ wrt } V$	med.regressCOP2	copula theory
Quantile regression	$V \text{ wrt } U$	qua.regressCOP	copula theory
Quantile regression	$U \text{ wrt } V$	qua.regressCOP2	copula theory
Copula section	$t \mapsto \mathbf{C}(t, a)$	sectionCOP	copula theory
Copula section	$t \mapsto \mathbf{C}(a, t)$	sectionCOP	copula theory

The previous three tables do not list all of the myriad of special functions to support similar operations on *empirical copulas*. All empirical copula operators and utilities are prepended with **EMPIR** in the function name. An additional note concerning package nomenclature is that an appended “2” to a function name indicates $U \text{ wrt } V$ (e.g. **EMPIRgridderinv2** for an inversion of the partial derivatives $\delta \mathbf{C} / \delta v$ across the grid of the empirical copula).

Some additional functions to compute often salient features or characteristics of copulas or bivariate data, including functions for bivariate inference or goodness-of-fit, are listed in the following table:

Name	Symbol	Function	Concept
Left-tail decreasing	$V \text{ wrt } U$	isCOP.LTD	bivariate association
Left-tail decreasing	$U \text{ wrt } V$	isCOP.LTD	bivariate association
Right-tail increasing	$V \text{ wrt } U$	isCOP.RTI	bivariate association
Right-tail increasing	$U \text{ wrt } V$	isCOP.RTI	bivariate association
Pseudo-polar representation	$(\widehat{S}, \widehat{W})$	psepolar	extremal dependency
Tail concentration function	$q_C(t)$	tailconCOP	bivariate tail association
Tail (lower) dependency	λ_C^L	taildepCOP	bivariate tail association
Tail (upper) dependency	λ_C^U	taildepCOP	bivariate tail association
Tail (lower) order	κ_C^L	tailordCOP	bivariate tail association
Tail (upper) order	κ_C^U	tailordCOP	bivariate tail association
Neg'yly quadrant dependency	NQD	isCOP.PQD	bivariate association
Pos'yly quadrant dependency	PQD	isCOP.PQD	bivariate association
Permutation symmetry	permsym	isCOP.permsym	copula symmetry
Radial symmetry	radsym	isCOP.radsym	copula symmetry
Skewness (Joe, 2014)	$\eta(p; \psi)$	uvskew	bivariate skewness
Kullback–Leibler Divergence	$KL(f \mid g)$	kullCOP	bivariate inference
KL sample size	n_{fg}	kullCOP	bivariate inference
The Vuong Procedure	—	vuongCOP	bivariate inference
Spectral measure	$H(w)$	spectralmeas	extremal dependency inference
Stable tail dependence	$\widehat{l}(x, y)$	stabtaildepf	extremal dependency inference
L-comoments (samp. distr.)	—	lcomCOPpv	experimental bivariate inference

The **Table of Probabilities** that follows lists important relations between various joint probability concepts, the copula, nonexceedance probabilities u and v , and exceedance probabilities u' and v' . A compact summary of these probability relations has obvious usefulness. The notation $[\dots, \dots]$ is to read as $[\dots \text{ and } \dots]$, and the $[\dots \mid \dots]$ is to be read as $[\dots \text{ given } \dots]$.

Probability	and	Symbol Convention
$\Pr[U \leq u, V \leq v]$	=	$\mathbf{C}(u, v)$ — The copula, COP
$\Pr[U > u, V > v]$	=	$\hat{\mathbf{C}}(u', v')$ — The survival copula, surCOP
$\Pr[U \leq u, V > v]$	=	$u - \mathbf{C}(u, v')$
$\Pr[U > u, V \leq v]$	=	$v - \mathbf{C}(u', v)$
$\Pr[U \leq u \mid V \leq v]$	=	$\mathbf{C}(u, v)/v$
$\Pr[V \leq v \mid U \leq u]$	=	$\mathbf{C}(u, v)/u$
$\Pr[U \leq u \mid V > v]$	=	$(u - \mathbf{C}(u, v))/(1 - v)$
$\Pr[V \leq v \mid U > u]$	=	$(v - \mathbf{C}(u, v))/(1 - u)$
$\Pr[U > u \mid V > v]$	=	$\hat{\mathbf{C}}(u', v')/u' = \overline{\mathbf{C}}(u, v)/(1 - u)$
$\Pr[V > v \mid U > u]$	=	$\hat{\mathbf{C}}(u', v')/v' = \overline{\mathbf{C}}(u, v)/(1 - v)$
$\Pr[V \leq v \mid U = u]$	=	$\delta \mathbf{C}(u, v)/\delta u$ — Partial derivative, derCOP
$\Pr[U \leq u \mid V = v]$	=	$\delta \mathbf{C}(u, v)/\delta v$ — Partial derivative, derCOP2
$\Pr[U > u \text{ or } V > v]$	=	$\mathbf{C}^*(u', v') = 1 - \mathbf{C}(u', v')$ — The co-copula, coCOP
$\Pr[U \leq u \text{ or } V \leq v]$	=	$\tilde{\mathbf{C}}(u, v) = u + v - \mathbf{C}(u, v)$ — The dual of a copula, duCOP

Table of Conditional Expectations that follows includes *conditional expectation* (*conditional mean*) ([EuvCOP](#), [EvuCOP](#)) and *conditional L-moments*:

Probability	and	Symbol Convention
$E[V U = u]$	=	$\int_0^1 (1 - \delta \mathbf{C}(u, v) / \delta u) dv$ — Expectation of V given U, EuvCOP
$E[U V = v]$	=	$\int_0^1 (1 - \delta \mathbf{C}(u, v) / \delta v) du$ — Expectation of U given V, EuvCOP
$(\lambda_r, \tau_r)[V U = u]$	$\mapsto$	uses derCOP — conditional L-moments of V given U, LMRvuCOP
$(\lambda_r, \tau_r)[U V = v]$	$\mapsto$	uses derCOP2 — conditional L-moments of U given V, LMRuvCOP

The function [jointCOP](#) has considerable demonstration in its **Note** section of the **joint and joint or** relations shown through simulation and counting scenarios. Also there is a demonstration in the **Note** section of function [duCOP](#) on application of the concepts of **joint and** conditions, **joint or** conditions, and importantly joint **mutually exclusive or** conditions.

Copula Construction Methods

Permutation asymmetry (nonexchangeability) can be added to (inserted in) a copula by [breveCOP](#). One, two, or more copulas can be “composited,” “combined,” or “multiplied” in interesting ways to create highly unique bivariate relations and as a result, complex dependence structures can be formed. Such operations can often insert permutation asymmetry as well. To such an end, the package provides three main functions for copula composition: [composite1COP](#) composites a single copula with two compositing parameters (*Khoudraji device* with *independence*), [composite2COP](#) (*Khoudraji device*) composites two copulas with two compositing parameters, and [composite3COP](#) composites two copulas with four compositing parameters.

Two copulas also can be combined through a weighted convex combination using [convex2COP](#) with a single weighting parameter, and even N number of copulas can be combined by weights using [convexCOP](#). So-called “gluing” two copula by a parameter is provided by [glueCOP](#). Multiplication of two copulas to form a third is supported by [prod2COP](#). All eight functions for compositing, combining, or multiplying copulas are compatible with joint probability simulation ([simCOP](#)), measures of association (e.g. $\rho_{\mathbf{C}}$), and presumably all other copula operations using **copBasic** features. Finally, *ordinal sums* of copula are provided by [ORDSUMcop](#) and [ORDSUWcop](#) as particularly interesting methods of combining copulas.

No. of copulas	Combining Parameters	Function	Concept
1	β	breveCOP	adding permutation asymmetry
1	α, β	composite1COP	copula combination
1	α, β	khoudraji1COP	copula combination
1	α, β	khoudrajiPCOP	copula combination
2	α, β	composite2COP	copula combination
2	α, β	khoudraji2COP	copula combination
2	$\alpha, \beta, \kappa, \gamma$	composite3COP	copula combination
2	$\alpha, (1 - \alpha)$	convex2COP	weighted copula combination
N	$\omega_{i \in N}$	convexCOP	weighted copula combination
2	γ	glueCOP	gluing of coupla
2	$(\mathbf{C}_1 * \mathbf{C}_2)$	prod2COP	copula multiplication
N	$\mathbf{C}_{\mathcal{J}_i}$ for $\mathcal{J}_{i \in N}$ partitions	ORDSUMcop	M -ordinal sums of copulas
N	$\mathbf{C}_{\mathcal{J}_i}$ for $\mathcal{J}_{i \in N}$ partitions	ORDSUWcop	W -ordinal sums of copulas

Useful Copula Relations by Visualization

There are a myriad of relations amongst variables computable through copulas, and these were listed in the **Table of Probabilities** earlier in this documentation. There is a script located in the `inst/doc` directory of the **copBasic** sources titled `CopulaRelations_BaseFigure_inR.txt`. This script demonstrates, using the **PSP** copula, relations between the copula (**COP**), survival copula (**surCOP**), joint survival function of a copula (**surfuncCOP**), co-copula (**coCOP**), and dual of a copula function (**duCOP**). The script performs simulation and manual counts observations meeting various criteria in order to compute their *empirical probabilities*. The script produces a base figure, which after extending in editing software, is suitable for educational description and is provided at the end of this documentation.

A Review of “Return Periods” using Copulas

Study of natural hazards commonly use as *annual return periods* T in years, which are defined for a nonexceedance probability q as $T = 1/(1 - q)$. In bivariate analysis, there immediately emerge two types of return periods representing $T_{q; \text{coop}}$ and $T_{q; \text{dual}}$ conditions between nonexceedances of the two hazard sources (random variables) U and V . It is usual in many applications for T to be expressed equivalently as a probability q in common for both variables.

Incidentally, the $\Pr[U > u \mid V > v]$ and $\Pr[V > v \mid U > u]$ probabilities also are useful for *conditional return period* computations following Salvadori *et al.* (2007, pp. 159–160) but are not further considered here. Also the $F_K(w)$ (*Kendall Function* or *Kendall Measure* of a copula) is the core tool for *secondary return period* computations (see **kfuncCOP**).

Let the copula $\mathbf{C}(u, v; \Theta)$ for nonexceedances u and v be set for some copula family (formula) by a parameter vector Θ . The copula family and parameters define the joint coupling (loosely meant the dependency/correlation) between hazards U and V . If “failure” occurs if **either** or **both** hazards U and V are at probability q threshold ($u = v = 1 - 1/T = q$) for T -year return period, then the **real return period** of failure is defined using either the copula $\mathbf{C}(q, q; \Theta)$ or the *co-copula* $\mathbf{C}^*(q', q'; \Theta)$ for exceedance probability $q' = 1 - q$ is

$$T_{q; \text{coop}} = \frac{1}{1 - \mathbf{C}(q, q; \Theta)} = \frac{1}{\mathbf{C}^*(1 - q, 1 - q; \Theta)} \text{ and}$$

$$T_{q; \text{coop}} \equiv \frac{1}{\text{cooperative risk}}.$$

Or in words, the hazard sources **collaborate** or **cooperate** to cause failure. If failure occurs, however, if and only if **both** hazards U and V occur simultaneously (the hazards must “dually work together” or be “conjunctive”), then the **real return period** is defined using either the *dual of a copula (function)* $\tilde{\mathbf{C}}(q, q; \Theta)$, the *joint survival function* $\overline{\mathbf{C}}(q, q; \Theta)$, or *survival copula* $\hat{\mathbf{C}}(q', q'; \Theta)$ as

$$T_{q; \text{dual}} = \frac{1}{1 - \tilde{\mathbf{C}}(q, q; \Theta)} = \frac{1}{\overline{\mathbf{C}}(q, q; \Theta)} = \frac{1}{\hat{\mathbf{C}}(q', q'; \Theta)} \text{ and}$$

$$T_{q; \text{dual}} \equiv \frac{1}{\text{complement of dual protection}}.$$

Numerical demonstration is informative. Salvadori *et al.* (2007, p. 151) show for a *Gumbel–Hougaard copula* (**GHcop**) having $\Theta = 3.055$ and $T = 1,000$ years ($q = 0.999$) that $T_{q; \text{coop}} = 797.1$ years and that $T_{q; \text{dual}} = 1,341.4$ years, which means that average return periods between “failures” are

$$T_{q; \text{coop}} \leq T \leq T_{q; \text{dual}} \text{ and thus}$$

$$797.1 \leq T \leq 1314.4 \text{ years.}$$

With the following code, these bounding return-period values are readily computed and verified using the `prob2T()` function from the **lmomco** package along with **copBasic** functions `COP` (generic functional interface to a copula) and `duCOP` (*dual of a copula*):

```
q <- lmomco::T2prob(1000)
lmomco::prob2T( COP(q,q, cop=GHcop, para=3.055)) # 797.110
lmomco::prob2T(duCOP(q,q, cop=GHcop, para=3.055)) # 1341.438
```

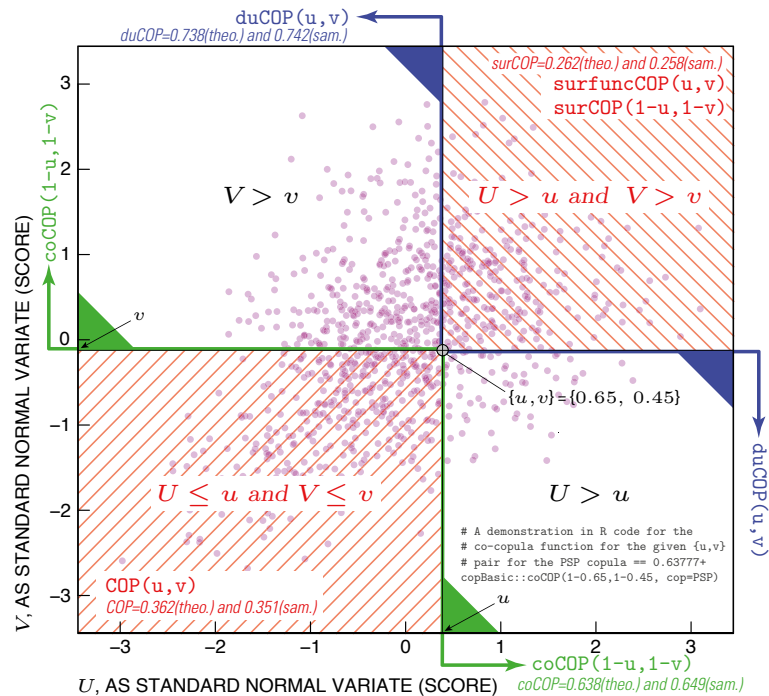
An early source (in 2005) by some of those authors cited on p. 151 of Salvadori *et al.* (2007; their citation “[67]”) shows $T_{q; \text{dual}} = 798$ years—a rounding error seems to have been committed. Finally and just for reference, a Gumbel–Hougaard copula having $\Theta = 3.055$ corresponds to an analytical *Kendall Tau* (see `GHcop`) of $\tau \approx 0.673$, which can be verified through numerical integration available from `tauCOP` as:

```
tauCOP(cop=GHcop, para=3.055, brute=TRUE) # 0.6726542
```

Thus, a “better understanding of the statistical characteristics of [multiple hazard sources] requires the study of their joint distribution” (Salvadori *et al.*, 2007, p. 150).

Interaction of copBasic to the Copulas and Features of Other Packages

The **copBasic** package (c.2008–11) was not originally intended to be a port of the numerous bivariate copulas or over reimplement other bivariate copulas available in **R**, but in support of didactic and teaching efforts. Though as the package passed its 10th year in 2018, the original intent changed to a more generalized purpose as more copula were added and other features by request. It is useful to point out a demonstration showing an implementation of the *Gaussian copula* from the **copula** package, which is shown in the **Note** section of `med.regressCOP` in a circumstance of ordinary least squares linear regression compared to *median regression* of a copula as well as prediction limits of both regressions. Another demonstration in context of *maximum pseudo-log-likelihood estimation* of copula parameters is seen in the **Note** section `mleCOP`, and also see “**API to the copula package**” or “**package copula (comparison to)**” entries in the Index of this user manual.



EXPLANATION

- Simulated value from parameterless "PSP" copula with a sample size of 1,000
- $duCOP(u, v)$ The monospace typeface indicates commands in R using copBasic syntax
- $surCOP=0.262(\text{theo.})$ and $0.258(\text{sam.})$ The slanted typeface indicates theoretical values from the copula and sample estimates from the data—For this example, the survival copula has probability $\Pr[U > u \text{ and } V > v] = 0.262$ and the simulations have 258 occurrences in the respectively shown hatched region.

Author(s)

William Asquith <william.asquith@ttu.edu>

References

- Cherubini, U., Luciano, E., and Vecchiato, W., 2004, Copula methods in finance: Hoboken, NJ, Wiley, 293 p.
- Hernández-Maldonado, V., Díaz-Viera, M., and Erdely, A., 2012, A joint stochastic simulation method using the Bernstein copula as a flexible tool for modeling nonlinear dependence structures between petrophysical properties: Journal of Petroleum Science and Engineering, v. 90–91, pp. 112–123.
- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
- Hofert, M., Kojadinovic, I., Mächler, M., and Yan, J., 2018, Elements of copula modeling with R: Dordrecht, Netherlands, Springer.
- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, Extremes in nature—An approach using copulas: Dordrecht, Netherlands, Springer, Water Science and Technology Library 56, 292 p.

Examples

```
## Not run:
# Nelsen (2006, p. 75, exer. 3.15b) provides for a nice test of copBasic features.
"mcdurv" <- function(u,v, theta) {
  ifelse(u > theta & u < 1-theta & v > theta & v < 1 - theta,
    return(M(u,v) - theta), # Upper bounds copula with a shift
    return(W(u,v)))        # Lower bounds copula
}
"MCDURV" <- function(u,v, para=NULL) {
  if(is.null(para)) stop("need theta")
  if(para < 0 | para > 0.5) stop("theta ! in [0, 1/2]")
  return(asCOP(u, v, f=mcdurv, para))
}
"afunc" <- function(t) { # a sample size = 1,000 hard wired
  return(cov(simCOP(n=1000, cop=MCDURV, para=t, ploton=FALSE, points=FALSE))[1,2])
}
set.seed(6234) # setup covariance based on parameter "t" and the "root" parameter
print(uniroot(afunc, c(0, 0.5))) # "t" by simulation = 0.1023742
# Nelsen reports that if theta approx. 0.103 then covariance of U and V is zero.
# So, one will have mutually completely dependent uncorrelated uniform variables!

# Let us check some familiar measures of association:
rhoCOP( cop=MCDURV, para=0.1023742) # Spearman Rho = 0.005854481 (near zero)
tauCOP( cop=MCDURV, para=0.1023742) # Kendall Tau = 0.2648521
wolfCOP(cop=MCDURV, para=0.1023742) # S & W Sigma = 0.4690174 (less familiar)
D <- simCOP(n=1000, cop=MCDURV, para=0.1023742) # Plot mimics Nelsen (2006, fig. 3.11)
# L-comoments (matrices) measure high dimensional variable comovements
# L-comoments (refer also to the lmomco package)
lcomCOP(cop=MCDURV, para=0.1023742)
lmomco::lcomoms2(simCOP(n=1000, cop=MCDURV, para=0.1023742), nmom=5)
lmomco::lcomoms2(simCOP(n=1000, cop=MCDURV, para=0), nmom=5) # Perfect neg. corr.
lmomco::lcomoms2(simCOP(n=1000, cop=MCDURV, para=0.5), nmom=5) # Perfect pos. corr.
# T2 (L-correlation), T3 (L-coskew), T4 (L-cokurtosis), and T5 matrices result. For
# Theta = 0 or 0.5 see the matrix symmetry with a sign change for L-coskew and T5 on
# the off diagonals (offdiags). See unities for T2. See near zero for offdiag terms
# in T2 near zero. But then see that T4 off diagonals are quite different from those
# for Theta 0.1024 relative to 0 or 0.5. As a result, T4 has captured a unique
# property of U vs V.
## End(Not run)
```

aicCOP

Akaike Information Criterion between a Fitted Coupla and an Empirical Copula

Description

Compute the *Akaike information criterion* (AIC) AIC_C (Chen and Guo, 2019, p. 29), which is computed using *mean square error* MSE_C as

$$\text{MSE}_C = \frac{1}{n} \sum_{i=1}^n (\mathbf{C}_n(u_i, v_i) - \mathbf{C}_{\Theta_m}(u_i, v_i))^2 \text{ and}$$

$$\text{AIC}_C = 2m + n \log(\text{MSE}_C),$$

where $\mathbf{C}_n(u_i, v_i)$ is the *empirical copula* (empirical joint probability) for the i th observation, $\mathbf{C}_{\Theta_m}(u_i, v_i)$ is the fitted copula having m parameters in Θ . The $\mathbf{C}_n(u_i, v_i)$ comes from [EMPIRCOP](#). The AIC_C is in effect saying that the best copula will have its joint probabilities plotting on a 1:1 line with the empirical joint probabilities, which is an $\text{AIC}_C = -\infty$. From the MSE_C shown above, the root mean square error [rmseCOP](#) and Bayesian information criterion (BIC) [bicCOP](#) can be computed. These goodness-of-fits can assist in deciding one copula favorability over another, and another goodness-of-fit using the absolute differences between $\mathbf{C}_n(u, v)$ and $\mathbf{C}_{\Theta_m}(u, v)$ is found under [statTn](#).

Usage

```
aicCOP(u, v=NULL, cop=NULL, para=NULL, m=NA, ...)
```

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction; If not given, then a second column from argument u is attempted;
cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
m	The number of parameters in the copula, which is usually determined by length of para if m=NA, but some complex compositions of copulas are difficult to authoritatively probe for total parameter lengths and mixing coefficients; and
...	Additional arguments to pass to either copula (likely most commonly to the empirical copula).

Value

The value for AIC_C is returned.

Author(s)

W.H. Asquith

References

Chen, Lu, and Guo, Shenglian, 2019, Copulas and its application in hydrology and water resources: Springer Nature, Singapore, ISBN 978–981–13–0574–0.

See Also

[EMPIRCOP](#), [bicCOP](#), [rmseCOP](#)

Examples

```
## Not run:
S <- simCOP(80, cop=GHcop, para=5) # Simulate some probabilities, but we
# must then treat these as data and recompute empirical probabilities.
U <- lmomco::pp(S$U, sort=FALSE); V <- lmomco::pp(S$V, sort=FALSE)
# The parent distribution is Gumbel-Hougaard extreme value copula, but in practical
# applications, we do not know that. Say we speculate that perhaps the Galambos extreme
# value might be the parent; maximum likelihood is used to fit the single parameter.
pGL <- mleCOP( U,V, cop=GLcop, interval=c(0, 20))$par
aics <- c(aicCOP(U,V, cop=GLcop, para=pGL), aicCOP(U,V, cop=P), aicCOP(U,V, cop=PSP))
names(aics) <- c("GLcop", "P", "PSP")
print(aics) # We will see that the first AIC is the smallest because the
# Galambos has the nearest overall behavior than the P and PSP copulas.
## End(Not run)
```

AMHcop

The Ali–Mikhail–Haq Copula

Description

The *Ali–Mikhail–Haq copula* (Nelsen, 2006, pp. 92–93, 172) is

$$C_{\Theta}(u, v) = \mathbf{AMH}(u, v) = \frac{uv}{1 - \Theta(1 - u)(1 - v)},$$

where $\Theta \in [-1, +1]$, where the right boundary, $\Theta = 1$, can sometimes be considered valid according to Mächler (2014). The copula $\Theta \rightarrow 0$ becomes the *independence copula* ($\Pi(u, v)$; [P](#)), and the parameter Θ is readily computed from a *Kendall Tau* ([tauCOP](#)) by

$$\tau_C = \frac{3\Theta - 2}{3\Theta} - \frac{2(1 - \Theta)^2 \log(1 - \Theta)}{3\Theta^2},$$

and by *Spearman Rho* ([rhoCOP](#)), through Mächler (2014), by

$$\rho_C = \sum_{k=1}^{\infty} \frac{3\Theta^k}{\binom{k+2}{2}^2}.$$

The support of τ_C is $[(5 - 8 \log(2))/3, 1/3] \approx [-0.1817258, 0.3333333]$ and the ρ_C is $[33 - 48 \log(2), 4\pi^2 - 39] \approx [-0.2710647, 0.4784176]$, which shows that this copula has a limited range of dependency. The infinite summation is easier to work with than Nelsen (2006, p. 172) definition of

$$\rho_C = \frac{12(1 + \Theta)}{\Theta^2} \text{dilog}(1 - \Theta) - \frac{24(1 - \Theta)}{\Theta^2} \log(1 - \Theta) - \frac{3(\Theta + 12)}{\Theta},$$

where the $\text{dilog}(x)$ is the dilogarithm function defined by

$$\text{dilog}(x) = \int_1^x \frac{\log(t)}{1 - t} dt.$$

The integral version has more nuances with approaches toward $\Theta = 0$ and $\Theta = 1$ than the infinite sum version.

Usage

```
AMHcop(u, v, para=NULL, rho=NULL, tau=NULL, fit=c("rho", "tau"), ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula. However, if a second parameter is present, it is treated as a logical to reverse the copula ($u + v - 1 + \mathbf{AMH}(1 - u, 1 - v; \Theta)$);
<code>rho</code>	Optional Spearman Rho from which the parameter will be estimated and presence of rho trumps tau;
<code>tau</code>	Optional Kendall Tau from which the parameter will be estimated;
<code>fit</code>	If para, rho, and tau are all NULL, the <code>u</code> and <code>v</code> represent the sample. The measure of association by the <code>fit</code> declaration will be computed and the parameter estimated subsequently. The <code>fit</code> has not other utility than to trigger which measure of association is computed internally by the <code>cor</code> function in R; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned. Otherwise if tau is given, then the Θ is computed and a list having

<code>para</code>	The parameter Θ , and
<code>tau</code>	Kendall Tau.

and if para=NULL and tau=NULL, then the values within `u` and `v` are used to compute Kendall Tau and then compute the parameter, and these are returned in the aforementioned list.

Note

Mächler (2014) reports on accurate computation of τ_C and ρ_C for this copula for conditions of $\Theta \rightarrow 0$ and in particular derives the following equation, which does not have Θ in the denominator:

$$\rho_C = \sum_{k=1}^{\infty} \frac{3\Theta^k}{\binom{k+2}{2}^2}.$$

The **copula** package provides a Taylor series expansion for τ_C for small Θ in the `copula::tauAMH()`. This is demonstrated here between the implementation of $\tau = 0$ for parameter estimation in the **copBasic** package to that in the more sophisticated implementation in the **copula** package.

```
copula::tauAMH(AMHcop(tau=0)$para) # theta = -2.313076e-07
```

It is seen that the numerical approaches yield quite similar results for small τ_C , and finally, a comparison to the ρ_C is informative:

```
rhoCOP(AMHcop, para=1E-9) # 3.333333e-10 (two nested integrations)
copula:::rhoAmhCopula(1E-9) # 3.333333e-10 (cutoff based)
theta <- seq(-1,1, by=.0001)
RHOa <- sapply(theta, function(t) rhoCOP(AMHcop, para=t))
RHOb <- sapply(theta, function(t) copula:::rhoAmhCopula(t))
plot(10^theta, RHOa-RHOb, type="l", col=2)
```

The plot shows that the apparent differences are less than 1 part in 100 million—The **copBasic** computation is radically slower though, but **rhoCOP** was designed for generality of copula family.

Author(s)

W.H. Asquith

References

- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
- Mächler, Martin, 2014, Spearman's Rho for the AMH copula—A beautiful formula: copula package vignette, accessed on April 7, 2018, at <https://CRAN.R-project.org/package=copula> under the vignette *rhoAMH-dilog.pdf*.
- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Pranesh, Kumar, 2010, Probability distributions and estimation of Ali–Mikhail–Haq copula: Applied Mathematical Sciences, v. 4, no. 14, p. 657–666.

See Also

[P](#)

Examples

```
## Not run:
t <- 0.9 # The Theta of the copula and we will compute Spearman Rho.
di <- integrate(function(t) log(t)/(1-t), lower=1, upper=(1-t))$value
A <- di*(1+t) - 2*log(1-t) + 2*t*log(1-t) - 3*t # Nelsen (2007, p. 172)
rho <- 12*A/t^2 - 3 # 0.4070369
rhoCOP(AMHcop, para=t) # 0.4070369
sum(sapply(1:100, function(k) 3*t^k/choose(k+2, 2)^2)) # Machler (2014)
# 0.4070369 (see Note section, very many tens of terms are needed)
## End(Not run)

## Not run:
layout(matrix(1:2, byrow=TRUE)) # Note: Kendall Tau is same on reversal.
s <- 2; set.seed(s); nsim <- 10000
UVn <- simCOP(nsim, cop=AMHcop, para=c(-0.9, "FALSE"), col=4)
mtext("Normal definition [default]") # '2nd' parameter could be skipped
set.seed(s) # seed used to keep Rho/Tau inside attainable limits
UVr <- simCOP(nsim, cop=AMHcop, para=c(-0.9, "TRUE"), col=2)
mtext("Reversed definition")
AMHcop(UVn[,1], UVn[,2], fit="rho")$rho # -0.2581653
AMHcop(UVr[,1], UVr[,2], fit="rho")$rho # -0.2570689
```

```

rhoCOP(cop=AMHcop, para=-0.9)          # -0.2483124
AMHcop(UVn[,1], UVn[,2], fit="tau")$tau # -0.1731904
AMHcop(UVr[,1], UVr[,2], fit="tau")$tau # -0.1724820
tauCOP(cop=AMHcop, para=-0.9)          # -0.1663313
## End(Not run)

```

asCOP

Wrapper on a User-Level Formula to Become a Copula Function

Description

This function is intended to document and then to extend a simple API to end users to aid in implementation of other copulas for use within the **copBasic** package. There is no need or requirement to use asCOP for almost all users. However, for the mathematical definition of some copulas, the asCOP function might help considerably. This is because there is a need for special treatment of u and v vectors of probability as each interacts with the vectorization implicit in R. The special treatment is needed because many copulas are based on the operators such as `min()` and `max()`. When numerical integration used by the `integrate()` function in R in some copula operators, such as `tauCOP` for the *Kendall Tau* of a copula, special accommodation is needed related to the inherent vectorization in R and how `integrate()` works.

Basically, the problem is that one can not strictly rely in all circumstances on what R does in terms of value recycling when u and v are of unequal lengths. The source code is straightforward. Simply put, if lengths of u and v are unity, then there is no concern, and even if the length of u (say) is unity and v is 21, then recycling of u would often be okay. The real danger is when u and v have unequal lengths and those lengths are each greater than unity—the R treatment can not be universally relied upon with the various numerics herein involving optimization and nested numerical integration.

The example shows how a formula definition of a copula that is not a copula already implemented by **copBasic** is set into a function `delTacop` and then used inside another function `UsersCop` that will be the official copula that is compatible with a host of functions in **copBasic**. The use of asCOP provides the length check necessary on u and v , and the argument `...` provides optional parameter support should the user's formula require more settings.

Usage

```
asCOP(u, v, f=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>f</code>	A function for which the user desires to make as a copula; and
<code>...</code>	Additional arguments to pass to the function <code>f</code> (such as parameters, if needed, for the copula in the form of a list).

Value

The value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also[COP](#)**Examples**

```
## Not run:
# Concerning Nelsen (2006, exer. 3.7, pp. 64--65)
"trianglecop" <- function(u,v, para=NULL, ...) {
  # If para is set, then the triangle is rotated 90d clockwise.
  if(! is.null(para) && para == 1) { t <- u; u <- v; v <- t }
  if(length(u) > 1 | length(v) > 1) stop("only scalars for this function")
  v2<-v/2; if(0 <= u & u <= v2 & v2 <= 1/2) { return(u ) }
  } else if(0 <= v2 & v2 < u & u < 1-v2) { return(v2 ) }
  } else if(1/2 <= 1-v2 & 1-v2 <= u & u <= 1 ) { return(u+v-1) }
  } else { stop("should not be here in logic") }
}
"UsersCop" <- function(u,v, ...) { asCOP(u,v, f=trianglecop, ...) }
n=20000; UV <- simCOP(n=n, cop=UsersCop)
# The a-d elements of the problem now follow:
# (a) Pr[V = 1 - |2*U -1|] = 1 and Cov(U,V) = 0; so that two random variables
# can be uncorrelated but each is perfectly predictable from the other
mean(UV$V - (1 - abs(2*UV$U -1))) # near zero; Nelsen says == 0
cov(UV$U, UV$V) # near zero; Nelsen says == 0

# (b) Cop(m,n) = Cop(n,m); so that two random variables can be identically
# distributed, uncorrelated, and not exchangeable
EMPIRcop(0.95,0.17, para=UV) # = A
EMPIRcop(0.17,0.95, para=UV) # = B; then A != B

# (c) Pr[V - U > 0] = 2/3; so that two random variables can be identically
# distributed, but their difference need not be symmetric about zero
tmp <- (UV$V - UV$U) > 0
length(tmp[tmp == TRUE])/n # about 2/3; Nelsen says == 2/3
# the prior two lines yield about 1/2 for independence copula P()

# (d) Pr[X + Y > 0] = 2/3; so that uniform random variables on (-1,1) can each
# be symmetric about zero, but their sum need not be.
tmp <- ((2*UV$V - 1) + (2*UV$U - 1)) > 0
length(tmp[tmp == TRUE])/n # about 2/3; Nelsen says == 2/3
## End(Not run)

## Not run:
# Concerning Nelsen (2006, exam. 3.10, p. 73)
"shufflecop" <- # assume scalar arguments for u and v
```

```

function(u,v, para, ...) {
  m <- para$mixer; subcop <- para$subcop
  if(is.na(m) | m <= 0 | m >= 1) stop("m ! in [0,1]")
  if(u <= m) { return(      subcop(1-m+u, v, para=para$para) -
                           subcop(1-m,   v, para=para$para))
  } else {      return(v - subcop(1-m,   v, para=para$para) +
                      subcop(u-m,   v, para=para$para))
  }
}
}
"UsersCop" <- function(u,v, para=NULL) {
  asCOP(u,v, f=shufflecop, para=para)
}
n <- 1000; u <- runif(n)
para <- list(mixer=runif(1), subcop=W, para=20)
v <- sapply(1:n, function(i) {
  simCOPmicro(u[i], cop=UsersCop, para=para) })
plot(data.frame(U=u, V=v), pch=17, col=rgb(1,0,1,1),
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
mtext("Shuffle Copula Nelsen (2006, exam. 3.10, p. 73)")

# Concerning Nelsen (2006, exam. 5.14, p. 195)
"deltacop" <- function(u,v, ...) { min(c(u,v,(u^2+v^2)/2))      }
"UsersCop" <- function(u,v, ...) { asCOP(u,v, f=deltacop, ...) }
isCOP.PQD(cop=UsersCop) # TRUE + Rho=0.288 and Tau=0.333 as Nelsen says
isCOP.LTD(cop=UsersCop, wrtV=TRUE) # FALSE as Nelsen says
isCOP.RTI(cop=UsersCop, wrtV=TRUE) # FALSE as Nelsen says
## End(Not run)

```

bicCOP

Bayesian Information Criterion between a Fitted Coupla and an Empirical Copula

Description

Compute the *Bayesian information criterion* (BIC) BIC_C (Chen and Guo, 2019, p. 29), which is computed using *mean square error* MSE_C as

$$MSE_C = \frac{1}{n} \sum_{i=1}^n (C_n(u_i, v_i) - C_{\Theta_m}(u_i, v_i))^2 \text{ and}$$

$$BIC_C = m \log(n) + n \log(MSE_C),$$

where $C_n(u_i, v_i)$ is the *empirical copula* (empirical joint probability) for the i th observation, $C_{\Theta_m}(u_i, v_i)$ is the fitted copula having m parameters in Θ . The $C_n(u_i, v_i)$ comes from [EMPIRCOP](#). The BIC_C is in effect saying that the best copula will have its joint probabilities plotting on a 1:1 line with the empirical joint probabilities, which is an $BIC_C = -\infty$. From the MSE_C shown above, the root mean square error [rmseCOP](#) and Akaike information criterion (AIC) [aicCOP](#) can be computed. These goodness-of-fits can assist in deciding on one copula favorability over another, and another goodness-of-fit using the absolute differences between $C_n(u, v)$ and $C_{\Theta_m}(u, v)$ is found under [statTn](#).

Usage

```
bicCOP(u, v=NULL, cop=NULL, para=NULL, m=NA, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction; If not given, then a second column from argument <code>u</code> is attempted;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>m</code>	The number of parameters in the copula, which is usually determined by length of <code>para</code> if <code>m=NA</code> , but some complex compositions of copulas are difficult to authoritatively probe for total parameter lengths and mixing coefficients; and
<code>...</code>	Additional arguments to pass to either copula (likely most commonly to the empirical copula).

Value

The value for BIC_G is returned.

Author(s)

W.H. Asquith

References

Chen, Lu, and Guo, Shenglian, 2019, Copulas and its application in hydrology and water resources: Springer Nature, Singapore, ISBN 978–981–13–0574–0.

See Also

[EMPIRCop](#), [aicCOP](#), [rmseCOP](#)

Examples

```
## Not run:
S <- simCOP(80, cop=GHcop, para=5) # Simulate some probabilities, but we
# must then treat these as data and recompute empirical probabilities.
U <- lmomco::pp(S$U, sort=FALSE); V <- lmomco::pp(S$V, sort=FALSE)
# The parent distribution is Gumbel-Hougaard extreme value copula.
# But in practical application we don't know that but say we speculate that
# perhaps the Galambos extreme value might be the parent. Then maximum
# likelihood is used on that copula to fit the single parameter.
pGL <- mleCOP(U,V, cop=GLcop, interval=c(0,20))$par

bics <- c(bicCOP(U,V, cop=GLcop, para=pGL), bicCOP(U,V, cop=P), bicCOP(U,V, cop=PSP))
names(bics) <- c("GLcop", "P", "PSP")
print(bics) # We will see that the first BIC is the smallest as the
# Galambos has the nearest overall behavior than the P and PSP copulas.
## End(Not run)
```

bicoploc

Analog to Line of Organic Correlation by Copula Diagonal

Description

HIGHLY EXPERIMENTAL AND SUBJECT TO OVERHAUL OR REMOVAL—Compute an analog to the *line of organic correlation (reduced major axis)* using the *diagonal* of a copula.

$$\Pr[U \leq u, V \leq v] = \mathbf{C}(u, v) = f.$$

The primary diagonal is defined as

$$\delta_{\mathbf{C}}(t) = \mathbf{C}(t, t) = f.$$

Two diagnostic plots can be plotted by the arguments available for this function. The plot for (U, V) coordinate nonexceedance probability domain along with the analyses involving the copula diagonal inversion comes first, which is followed by that for (X, Y) coordinate domain along with the well-known line of organic correlation by the method of L-moments and such a “line” (could be a curve) by copula diagonal inversion.

This much infrastructure written for flexibility in how a copula would interact for the purpose of estimation with moment preservation. The simple $u = v$ might be sufficient but let us have some flexibility.

Usage

```
bicoploc(xp, yp=NULL, xout=NA, xpara=NULL, ypara=NULL, dtypex="nor", dtypey="nor",
  ctype=c("weibull", "hazen", "bernstein", "checkerboard"), kumaraswamy=TRUE,
  plotuv=TRUE, plotxy=TRUE, adduv=FALSE, addxy=FALSE, snv=FALSE, limout=TRUE,
  autoleg=TRUE, xleg="topleft", yleg=NULL, rugxy=TRUE, ruglwd=0.5,
  xlim=NULL, ylim=NULL, titleuv="", titlexy="", titlecex=1,
  a=0, ff=pnorm(seq(-5, +5, by=0.1)), locdigits=6,
  paracop=TRUE, verbose=TRUE, x=NULL, y=NULL, ...)
```

Arguments

xp	Numeric vector giving paired data points of X . If this is a matrix or data frame, then the first and second columns are extracted for the xp and yp internal;
yp	Optional numeric vector giving paired data points of Y depending on the composition of x;
xout	An optional set of numeric values specifying where interpolation through the diagonal inversion is to take place;
xpara	An lmomco package parameter object for the X variable, which if not provided will trigger an method of L-moment parameter estimation for the distribution dtypex;

ypara	An lmomco package parameter object for the Y variable, which if not provided will trigger an method of L-moment parameter estimation for the distribution dtypey;
dtypex	The lmomco package distribution abbreviations (see <code>lmomco::dist.list()</code>) for the X variable. If this argument is set NULL and xpara is given with an element of type, then that distribution type is assigned internally to dtypex;
dtypey	The lmomco package distribution abbreviations (see <code>lmomco::dist.list()</code>) for the Y variable; If this argument is set NULL and xpara is given with an element of type, then that distribution type is assigned internally to dtypex;
ctype	Argument of the same name for the empirical copula for dispatch to EMPIRCop . The $1/n$ form is disabled for bicoploc operations based on limited experiments. The first letter of the argument's value is extracted, converted to upper case, and used as the plotting character in the two diagnostic plots;
kumaraswamy	A logical to trigger Kumaraswamy distribution smoothing of the copula diagonal inversion from the <i>empirical copula</i> . The Kumaraswamy distribution is a distribution having support $[0, 1]$ with an explicit quantile function and takes the place of a Beta distribution (see lmomco function <code>quakur()</code> for more details). The smoothing by Kumaraswamy will provide a continuous real number on the interval, which should insure no flat-lining as one rolls on to or off off the discrete interval provided by the empirical copula for expected sample sizes of the operations anticipated for the bicoploc function;
plotuv	A logical to trigger plotting of the analyses in the (U, V) coordinate nonexceedance probability domain along with the analyses involving the copula diagonal inversion. If set true, then adduv is set false internally;
plotxy	A logical to trigger plotting of the analyses in the (X, Y) coordinate domain along with the well-known line of organic correlation by the method of L-moments and such a "line" (could be a curve) by copula diagonal inversion. If set true, then addxy is set false internally;
adduv	A logical when set true will not call the <code>plot()</code> function for (U, V) coordinate nonexceedance probability domain but the other graphical operations of lines and points will be called;
addxy	A logical when set true will not call the <code>plot()</code> function for (X, Y) coordinate domain but the other graphical operations of lines and points will be called;
snv	A logical when set true will plot the (U, V) coordinate nonexceedance probability domain in units of standard normal variates;
limout	A logical when set true will plot the (X, Y) coordinate domain with horizontal and vertical axis limits inflated to the x_{out} and Y predictions;
autoleg	A logical when set will draw a legend for the plots if the plots are requested;
xleg	The value to become the argument <code>x</code> in the <code>legend()</code> call. The default setting is based on general assumption that this bicoploc() function is to be more commonly used in positive association circumstances between X and Y (positive Spearman Rho);
yleg	The value to become the argument <code>y</code> in the <code>legend()</code> call;

rugxy	Call rug() plotting operations on the X and Y values used in the parameter estimation of the parametric marginal distributions using the <code>xp</code> and <code>yp</code> , unless either or both have been overridden for the contents in <code>x</code> and (or) <code>y</code> arguments;
ruglwd	The line wide passed into rug(). Because plotting of small and large sample sizes can make it difficult in the smaller samples to see the line, it is judged useful to explicitly have this setting as an declared argument;
xlim	A numeric vector that if precisely of length 2 and no missing values therein, will override the horizontal limits of the plotxy plot of the (X, Y) domain. Otherwise, the contents of <code>xlim</code> , if not null, are inserted into a range computation for the limits to apply;
ylim	A numeric vector that if precisely of length 2 and no missing values therein, will override the vertical limits of the plotxy plot of the (X, Y) domain. Otherwise, the contents of <code>ylim</code> , if not null, are inserted into a range computation for the limits to apply;
titleuv	An optional title for the (U, V) domain plot;
titlexy	An optional title for the (X, Y) domain plot;
titlecex	The character expansion factor for the titles;
a	Value for the plotting-position formula for <code>lmomco::pp()</code> , default is <code>a=0</code> , which specifies <i>Weibull plotting positions</i> ;
ff	The nonexceedance joint probability of of the copula diagonal from which inversion computes the marginal nonexceedance probability values for $u = v = t$ as $C(t, t) = f$ where <code>ff</code> is the variable notation for the joint probability f ;
locdigits	Number of digits for rounding exclusive to the <code>loc</code> data frame produced in the returned list. The reasoning for this setting is that the expected application in practical circumstances will have discipline knowledge of the rounding depth suitable;
paracop	A logical triggering the use of the parametric asymmetric copula fit by numerical optimization to the (U, V) domain of the data (see Details);
verbose	Show messages of incremental progress with a incremental counter on the message;
x	Numeric vector of X values to be used in parameter estimation of the marginal parametric distribution if <code>xpara=NULL</code> and these values are internally replaced with <code>xp</code> (paired X) if not otherwise specified. This provides the ability to insert an alternative and presumably longer vector of the entire X sample without the restriction of individual values paired to the Y . A feature unique to providing the <code>x</code> is that missing values can be and are removed on the fly prior to parameter estimation of the marginal distribution. The <code>x</code> can be specific independent of <code>y</code> or even at all for either. Absolutely no provision is made that <code>x</code> can be a matrix or data frame holding the <code>y</code> ;
y	Numeric vector of Y values to be used in parameter estimation of the marginal parametric distribution if <code>ypara=NULL</code> and these values are internally replaced with <code>yp</code> (paired Y) if not otherwise specified. This provides the ability to insert an alternative and presumably longer vector of the entire Y sample without the restriction of individual values paired to the X . A feature unique to providing the <code>y</code> is that missing values can be and are removed on the fly prior to parameter

estimation of the marginal distribution. The x can be specific independent of y or even at all for either; and

... Additional arguments to pass.

Details

ON THE USE OF AN PARAMETRIC ASYMMETRIC COPULA—

Value

Lists, vectors, and data frames for the computations and predictions are returned.

organic	A list containing a data frame of the predictions for x_{out} by the conventional LOC (<code>locpair</code>) (see also <code>locsols\$lmrloc</code>), the estimates by the L-moments of the parameters of the marginal distributions (<code>locpara</code>), the estimates by copula diagonal with Kumaraswamy smoothing (if requested) (<code>bicoploc</code>), and the predictions based solely on the empirical copula approximation for the diagonal (<code>bicoploc_emp</code>). The <code>bicoploc</code> and <code>bicoploc_emp</code> are equal to each other if Kumaraswamy was not used. The <code>bicoploc</code> is intended to be the official output from the <code>bicoploc()</code> function. The furthest right column is <code>bicoploc_cop</code> and represents the predicting values using a parametric copula as fit to the (U, V) domain mapped into the (X, Y) domain as explained elsewhere in this documentation or sources;
locsols	A list of solutions to the LOC based (1) (<code>locpair</code>) on the conventional definition on the paired data (x_p and y_p) with the finiteness check previously described by <code>lmomco::lmrloc()</code> and (2) (<code>locpara</code>) the LOC solution <i>not on the paired data</i> but extractable from the L-moments of the parameters for the marginal distributions. Certain permutations of available features will either have the two L-moment solutions equal, or just the slopes equal, or differing in both intercept and slope. The <code>lmrloc</code> list contains both L-moment and product moment estimation of the LOC to adhere precisely to <code>lmomco::lmrloc()</code> output;
xpara	The parameters of the marginal distribution in X either as given in <code>xpara</code> , as estimated from x_p , or estimated from x by the method of L-moments through the lmomco package;
ypara	The parameters of the marginal distribution in Y either as given in <code>ypara</code> , as estimated from y_p , or estimated from y by the method of L-moments through the lmomco package;
faqs	A named vector containing some numerical facts about the operations and principally the requisite sample sizes involved are reported here;
faqscop	A named vector containing some numerical facts about the operations involving the fitting of the parametric asymmetric copula (Plackett by default) to the (U, V) domain; and
diag	A data frame containing information on the copula diagonal including the joint probability column <code>jtprob</code> (the ff as stand in for $C(t, t) = f$), the $u = v = t$ in column <code>uv</code> by the Kumaraswamy smooth (if requested), and the solely empirical copula version in column <code>uv_emp</code> . If the Kumaraswamy smooth is not used, then <code>uv</code> and <code>uv_emp</code> will be equal to each other. The furthest right column is <code>uv_cop</code>

and represents the values using a parametric copula as fit to the (U, V) domain as explained elsewhere in this documentation or sources.

Note

The use of a copula diagonal inversion for purposes of a line of organic correlation analog within the text-book literature and elsewhere is unknown to the developers (December 2023).

Though bicoploc has extensive logic for working through the copula diagonal, if we know the parent distribution and we just have the marginal probabilities equal to each other (the diagonal), then we recover the moments of Y simply with $u = v$:

```
library(lmomco)
ff    <- c(0.0001, seq(0.001, 0.999, by=0.001), 0.9999)
xpara <- lmomco::vec2par(c(3, 0.6, -0.4), type="pe3")
ypara <- lmomco::vec2par(c(3, 0.4, +0.6), type="pe3")
xx <- rlmomco(100000, xpara)
yy <- approx(qlmomco(ff, xpara), qlmomco(ff, ypara), xout=xx)$y
lmr2par(yy, type="aep4")$para
#      mu      sigma      gamma
# 2.9972742 0.3993914 0.5980866
```

Author(s)

W.H. Asquith

References

Kruskal, W.H., 1953, On the uniqueness of the line of organic correlation: Biometrics, vol. 9, no. 1, pp. 47–58, [doi:10.2307/3001632](https://doi.org/10.2307/3001632).

See Also

[diagCOPatf](#)

Examples

```
# paracop set to FALSE in these examples for speed
set.seed(4); nsim <- 50
X <- rnorm(nsim, mean=3, sd=0.6)
Y <- rnorm(nsim, mean=0, sd=0.2)
zz <- bicoploc(X,Y, xout=c(2.5, 3.5, 4), dtypex="nor", dtypey="nor", paracop=FALSE)
# cor(X,Y, method="spearman") # +0.0785114 POSITIVE

set.seed(1); nsim <- 50
X <- rnorm(nsim, mean=3, sd=0.6)
Y <- rnorm(nsim, mean=0, sd=0.2)
zz <- bicoploc(X,Y, xout=c(2.5, 3.5, 4), dtypex="nor", dtypey="nor", paracop=FALSE)
# cor(X,Y, method="spearman") # -0.1351741 NEGATIVE

set.seed(1); nsim <- 50
X <- rnorm(nsim, mean=3, sd=0.6)
```

```

Y <- 0.843*X + rnorm(nsim, mean=0, sd=0.2)
zz <- bicoploc(X,Y, xout=c(2.5, 3.5, 4), dtypex="nor", dtypey="nor", paracop=FALSE)
# cor(X,Y, method="spearman") # for nsim=1E6 RHO=0.92367

set.seed(1); nsim <- 50
X <- lmomco::rlmomco(nsim, lmomco::vec2par(c(3, 0.6, +0.5), type="pe3"))
Y <- 0.3*X^2 + lmomco::rlmomco(nsim, lmomco::vec2par(c(0, 0.4, 0), type="pe3"))
zz <- bicoploc(X,Y, xout=c(2.5, 3.5, 4.5), dtypex="nor", dtypey="nor", paracop=FALSE)
# cor(X,Y, method="spearman") # for nsim=1E6 RHO=0.92366

set.seed(1); nsim <- 50
X <- lmomco::rlmomco(nsim, lmomco::vec2par(c(3, 0.6, +0.5), type="pe3"))
Y <- 0.3*X^2 + lmomco::rlmomco(nsim, lmomco::vec2par(c(0, 0.4, 0), type="pe3"))
zz <- bicoploc(X,Y, xout=c(2.5, 3.5, 4.5), dtypex="gev", dtypey="gev", paracop=FALSE)

## Not run:
#####
# Image 800 samples in X and Y and though created as pairs, let us assume only
# 50 are actually paired for purposes of demonstration of specified parameters
# and (or) alternative x and y vectors providing the larger sample.
set.seed(1); nsim <- 800; npair <- 50
X <- lmomco::rlmomco(nsim, lmomco::vec2par(c(3, 0.6, +0.5), type="pe3"))
Y <- 0.3*X^2 + rnorm(nsim, mean=0, sd=0.3)
ix <- sample(seq_len(nsim), npair, replace=FALSE)
Xp <- X[ix]; Yp <- Y[ix]; dtypex <- "gev"; dtypey <- "gev"
xpara <- lmomco::lmr2par(X, type=dtypex); ypara <- lmomco::lmr2par(Y, type=dtypey)

# The next two bicoploc() calls produce identical results except for the density
# of data points along the axes for the rug plots. Ultimately, the same parameter
# estimates for the margins exists for both calls. The plotuv is disabled so that
# the user can tab between the two plotxy plots and see that they are the same.
zz <- bicoploc(Xp,Yp, xout=c(2.5, 3.5, 4.5), ruglwd=0.9, plotuv=FALSE,
               xpara=xpara, ypara=ypara, dtypex=NULL, dtypey=NULL)
mtext("Example of specific xpara and ypara from larger sample", line=0.5)

zz <- bicoploc(Xp,Yp, xout=c(2.5, 3.5, 4.5), ruglwd=0.9, plotuv=FALSE,
               xpara=xpara, ypara=ypara, dtypex=NULL, dtypey=NULL, x=X, y=Y)
mtext("Example of specific xpara and ypara from larger sample", line=0.5) #
## End(Not run)

## Not run:
#####
set.seed(1); nsim <- 50
UV <- rCOP(nsim, cop=breveCOP, para=list(cop=W, alpha=0, beta=0))
X <- qnorm(UV[,1], mean=3, sd=0.6)
Y <- qnorm(UV[,2], mean=2, sd=0.2)
zz <- bicoploc(X,Y, xout=c(1.5, 2.5, 3.5, 4), dtypex="nor", dtypey="nor")

set.seed(1); nsim <- 50
UV <- rCOP(nsim, cop=breveCOP, para=list(cop=W, alpha=0, beta=0.5))
X <- qnorm(UV[,1], mean=3, sd=0.6)
Y <- qnorm(UV[,2], mean=2, sd=0.2)
zz <- bicoploc(X,Y, xout=c(1.5, 2.5, 3.5, 4), dtypex="nor", dtypey="nor")

```

```

set.seed(1); nsim <- 50
UV <- rCOP(nsim, cop=breveCOP, para=list(cop=M_N5p12b, para=3, alpha=0, beta=0.5))
X <- qnorm(UV[,1], mean=3, sd=0.6)^2
Y <- qnorm(UV[,2], mean=2, sd=0.2)
zz <- bicoploc(X,Y, xout=c(1.5, 2.5, 3.5, 4), ylim=c(1,2.5),
               xleg="bottomleft", dtypex="aep4", dtypey="nor")

# A TRUE COUNTER EXAMPLE? Are the 1-v operations not sufficient depending on
# rotation? Is the secondary diagonal of a copula useful? Is there something wrong
# with the reflection operations as implemented at the end of November 2023?
# Should negatives be turned into positives by reversing Y within the internal logic?
# The solution current at end of November 2023 seems proper.
set.seed(1); nsim <- 500
para <- list(cop1=W, para1=4, cop2=GHcop, para2=c(30,.6, .9), alpha=0, beta=0.1)
UV <- rCOP(nsim, cop=composite2COP, para=para)
X <- -qexp(UV[,1], rate=10)+0.1 # Or the problem is a reflected exponential but the
Y <- qnorm(UV[,2], mean=2, sd=0.2) # exp version in lmomco can not handle
zz <- bicoploc(X,Y, xout=c(-0.05, 0, 0.2), dtypex="exp", dtypey="nor")
# Possibly, try another distribution.
zz <- bicoploc(X,Y, xout=c(-0.3, -0.2, 0), ylim=c(1,4), dtypex="aep4", dtypey="nor") #
## End(Not run)

## Not run:
#####
set.seed(1); nsim <- 200; npair <- 30
para <- list(cop=PLcop, para=80, alpha=0.3, beta=0.05)
UV <- rCOP(nsim, cop=composite1COP, para=para, resamv01=TRUE)
X <- lmomco::qlmomco(UV[,1], lmomco::vec2par(c(3, 0.6, +0.4), type="pe3"))
Y <- lmomco::qlmomco(UV[,2], lmomco::vec2par(c(3, 0.4, +0.0), type="pe3"))
ix <- sample(seq_len(nsim), npair, replace=FALSE)
Xp <- X[ix]; Yp <- Y[ix]; dtypex <- "pe3"; dtypey <- "pe3"
xpara <- lmomco::lmr2par(X, type=dtypex); ypara <- lmomco::lmr2par(Y, type=dtypey)
plot(10^X, 10^Y, log="xy", las=1, pch=21, lwd=0.8, col="black", bg="white",
     xlab="SOME RISK PHENOMENON IN X-DIRECTION",
     ylab="SOME RISK PHENOMENON IN Y-DIRECTION")
xout <- c(1.5, 2.5, 3.5, 4)
xlim <- c(1.5, 4.5); ylim <- c(1.8, 5.0)
zz <- bicoploc(Xp,Yp, xout=xout,xpara=xpara,ypara=ypara, xlim=xlim,ylim=ylim)

zz <- bicoploc(Xp,Yp, xout=xout,xpara=xpara,ypara=ypara, xlim=xlim,ylim=ylim,x=X,y=Y)
zz <- bicoploc(Xp,Yp, xout=xout,dtypex="pe3",dtypey="pe3",xlim=xlim,ylim=ylim,x=X,y=Y)

zz <- bicoploc(X, Y, xout=xout,dtypex="pe3",dtypey="pe3",xlim=xlim,ylim=ylim)#
## End(Not run)

```

Description

Attention: This function is deprecated in favor of `lcomCOP`, which uses only direct numerical `integrate()` on the integrals shown below. The `bilmoms` function is strictly based on Monte Carlo integration.

Compute the *bivariate L-moments (ratios)* $(\delta_{k;\mathbf{C}}^{[\dots]})$ of a copula $\mathbf{C}(u, v; \Theta)$ and remap these into the *L-comoment* matrix counterparts (Serfling and Xiao, 2007; Asquith, 2011) including *L-correlation* (*Spearman Rho*), *L-coskew*, and *L-cokurtosis*. As described by Brahimi *et al.* (2015), the first four bivariate L-moments $\delta_k^{[12]}$ for random variable $X^{(1)}$ or U with respect to (*wrt*) random variable $X^{(2)}$ or V are defined as

$$\delta_{1;\mathbf{C}}^{[12]} = 2 \int \int_{T^2} \mathbf{C}(u, v) \, du \, dv - \frac{1}{2},$$

$$\delta_{2;\mathbf{C}}^{[12]} = \int \int_{T^2} (12v - 6) \mathbf{C}(u, v) \, du \, dv - \frac{1}{2},$$

$$\delta_{3;\mathbf{C}}^{[12]} = \int \int_{T^2} (60v^2 - 60v + 12) \mathbf{C}(u, v) \, du \, dv - \frac{1}{2}, \text{ and}$$

$$\delta_{4;\mathbf{C}}^{[12]} = \int \int_{T^2} (280v^3 - 420v^2 + 180v - 20) \mathbf{C}(u, v) \, du \, dv - \frac{1}{2},$$

where the bivariate L-moments are related to the L-comoment ratios by

$$6\delta_k^{[12]} = \tau_{k+1}^{[12]} \quad \text{and} \quad 6\delta_k^{[21]} = \tau_{k+1}^{[21]},$$

where in other words, “the third bivariate L-moment $\delta_3^{[12]}$ is one sixth the L-cokurtosis $\tau_4^{[12]}$.” The first four bivariate L-moments yield the first five L-comoments (there is no first order L-comoment ratio). The terms and nomenclature are not easy and also the English grammar adjective “ratios” is not always consistent in the literature. The $\delta_{k;\mathbf{C}}^{[\dots]}$ are **ratios**, and the returned `bilmoms` element by this function holds matrices for the marginal means, marginal L-scales and L-coscales, and then the **ratio** L-comoments.

Similarly, the $\delta_k^{[21]}$ are computed by switching $u \rightarrow v$ in the polynomials within the above integrals multiplied to the copula in the system of equations with u . In general, $\delta_k^{[12]} \neq \delta_k^{[21]}$ for $k > 1$ unless in the case of *permutation symmetric* (`isCOP.permsym`) copulas. By theory, $\delta_1^{[12]} = \delta_1^{[21]} = \rho_{\mathbf{C}}/6$ where $\rho_{\mathbf{C}}$ is a *Spearman Rho* `rhoCOP`.

The integral for $\delta_{4;\mathbf{C}}^{[12]}$ does not appear in Brahimi *et al.* (2015) but this and the other forms are verified in the **Examples** and discussion in **Note**. The four $k \in [1, 2, 3, 4]$ for U *wrt* V and V *wrt* U comprise a full spectrum of system of seven (not eight) equations. One equation is lost because $\delta_1^{[12]} = \delta_1^{[21]}$.

Usage

```
bilmoms(cop=NULL, para=NULL, only.bilmoms=FALSE,
        n=1E5, halton=TRUE, usetime=TRUE, ...)
```

Arguments

<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>only.bilmoms</code>	A logical to trigger return of the δ_k and skip L-comoment computation;
<code>n</code>	The Monte Carlo integration size. The default seems to be at least an order of magnitude greater than needed for many applied problems;
<code>halton</code>	A logical triggering <i>Halton sequences</i> for the Monte Carlo integration instead of the bivariate uniform distribution (independence). The Sobol sequences are dependent on the <code>halton()</code> function of the randtoolbox package, and the Halton sequences canvas the $\mathcal{I}^2$ domain for smaller n values than required if statistical independence is used for the Monte Carlo integration. Note, randtoolbox at least at version 2.0.+ has “scrambling” of Sobol sequences temporarily disabled; therefore, Halton sequences with <code>usetime</code> set true is used herein (Spring 2026);
<code>usetime</code>	The argument of the same name for <code>randtoolbox::halton()</code> ; and
<code>...</code>	Additional arguments to pass to the <code>COP</code> and <code>rhoCOP</code> functions.

Value

An R list of the bivariate L-moments is returned.

<code>bilmomUV</code>	The bivariate L-moments $\delta_k^{[12]}$ of U with respect to V for $k \in [1, 2, 3, 4]$;
<code>bilmomVU</code>	The bivariate L-moments $\delta_k^{[21]}$ of V with respect to U for $k \in [1, 2, 3, 4]$;
<code>error.rho</code>	An “error” term in units of $\delta_1^{[12\&21]}$ used to judge whether the sample size for the Monte Carlo integration is sufficient based on a comparison to the <i>Spearman Rho</i> from direct numerical integration (not Monte Carlo based) using <code>rhoCOP</code> of the copula. Values for <code>error.rho</code> $< 1E-5$ seem to be sufficient to judge whether n is large enough;
<code>bilcomoms</code>	If not <code>only.bilmoms</code> , another R list holding the L-comoments (see Note) computed by simple remapping of the $\delta_k^{[\dots]}$ and parallel in structure to the function <code>lcomoms2()</code> of the lmomco package; and
<code>source</code>	An attribute identifying the computational source of the bivariate L-moments and bivariate L-comoments: “bilmoms.”

Note

The mapping of the bivariate L-moments to their L-comoment matrix counterparts is simple but nuances should be discussed and the meaning of the `error.rho` needs further description. The extra effort to form L-comoment matrices (Serfling and Xiao, 2007; Asquith, 2011) is made so that output matches the structure of the sample L-comoment matrices from the `lcomoms2()` function of the **lmomco** package.

Concerning the triangular or tent-shaped copula of Nelsen (2006, exer. 3.7, pp. 64–65) for demonstration, simulate from the triangular copula a sample of size $m = 20,000$ and compute some sample L-comoments using the following CPU intensive code. The function `asCOP` completes the vectorization needed for non-Monte Carlo integration for `rhoCOP`.

```

"trianglecop" <- function(u,v, para=NULL, ...) {
  # If para is set, then the triangle is rotated 90d clockwise.
  if(! is.null(para) && para == 1) { t <- u; u <- v; v <- t }
  if(length(u) > 1 | length(v) > 1) stop("only scalars for this function")
  v2<-v/2; if(0 <= u & u <= v2 & v2 <= 1/2) { return(u ) }
  } else if(0 <= v2 & v2 < u & u < 1-v2) { return(v2 ) }
  } else if(1/2 <= 1-v2 & 1-v2 <= u & u <= 1 ) { return(u+v-1) }
  } else { stop("should not be here in logic") }
}
"TriCop" <- function(u,v, ...) { asCOP(u,v, f=trianglecop, ...) }
m=20000; SampleUV <- simCOP(n=m, cop=TriCop, graphics=FALSE)
samLC <- lmomco::lcomoms2(SampleUV, nmom=5) # sample L-comoments
theoLC <- bilmoms(cop=TriCop)

```

The Error in Rho Computation—The ρ_C of the copula by numerical integration is computed internally to bilmoms as

```
rhoC <- rhoCOP(cop=TriCop) # -1.733858e-17
```

and used to compute the error.rho for bilmoms (see next code snippet). The ρ_C is obviously zero for this copula. Therefore, the bivariate association of TriCop is zero and thus is an example of a perfectly dependent situation yet of zero correlation. The bivariate L-moments and L-comoments of this copula are computed as

```
mean(replicate(20, bilmoms(cop=TriCop)$error.rho)) # 7.650723e-06
```

where the error.rho is repeated trials appears firmly $<1e-5$, which is near zero ($\epsilon_\rho \approx 0$). The error.rho term is defined by taking the first bivariate L-moment and numerically integrated ρ_C through `rhoCOP` and computing the terms

$$\begin{aligned}\epsilon_\rho^{[12]} &= |\delta_{1;C}^{[12]} - (\rho_C/6)|, \\ \epsilon_\rho^{[21]} &= |\delta_{1;C}^{[21]} - (\rho_C/6)|, \text{ and} \\ \epsilon_\rho &= \frac{\epsilon_\rho^{[12]} + \epsilon_\rho^{[21]}}{2},\end{aligned}$$

where the error.rho = ϵ_ρ , and values near zero are obviously favorable because this indicates that the Monte Carlo integration sample size n argument is sufficiently large to effectively canvas the $\mathcal{I}^2$ domain. For the situation here, the theoretical $\rho_C = 0$, but for $n = n = 100$, the error.rho ≈ 0.006 (e.g. `bilmoms(n=100, cop=TriCop)$error.rho`) through a 20-unit replication, which is a hint that 100 samples are not large enough and that should be obvious.

The reasoning behind using the error.rho between conventional numerical integration and the Monte Carlo integration (error.rho) is that ρ_C is symmetrical. This choice of “convergence” assessment reduces somewhat the sample size needed for Monte Carlo integration into single number representing error.

Discussion of Theoretical L-comoments—The theoretical L-comoments in the format structure of the sample L-comoments by the `lcomoms2()` function of the **lmomco** package are formed by the `bilmoms` function, and the theoretical values are shown below in sequence with details listed by L-comoment. Now we extract the L-comoment matrices and show the first L-comoment matrix (the matrix of the means):

```

theoLC1cm <- theoLC$bilmoms
print(theoLC1cm$L1)
      [,1]      [,2]
[1,] 0.4999939      NA
[2,]      NA 0.5000032

```

where the diagonal should be filled with 1/2, if the n is suitably large, because 1/2 is the mean of the marginal uniform random variables. By definition the secondary diagonal has NAs. The values shown above are extremely close supporting the idea that default n is large enough. The matrix of means is otherwise uninformative.

The second L-comoment matrix (L-scales and L-coscales) is

```

print(theoLC1cm$L2)
      [,1]      [,2]
[1,] 1.666677e-01 -3.466202e-06
[2,] -3.466209e-06 1.666674e-01

```

where the diagonal should be filled with 1/6, if the n is suitably large, because 1/6 is the univariate L-scale of the marginal uniform random variables. These values further support that default n is large enough. The diagonal is computed from the univariate L-moments of the margins of the Monte Carlo-generated edges and is otherwise uninformative. The secondary diagonal is a rescaling of the $\delta_1^{[\dots]}$ by the univariate L-moments of the margins to form L-coscales (nonratios). The copula is perfectly dependent but uncorrelated; so the secondary diagonal has near zeros.

The second L-comoment ratio matrix (coefficient of L-variations and L-correlations) is

```

print(theoLC1cm$T2)
      [,1]      [,2]
[1,] 1.000000e+00 -2.079712e-05
[2,] -2.079712e-05 1.000000e+00

```

where the diagonal by definition has unities (correlation is unity for a variable on itself) but the secondary diagonal for the L-correlations has near zeros because again the copula is uncorrelated, and the secondary diagonal is computed from the $\delta_1^{[\dots]}$. These L-correlations are the *Spearman Rho* values computed external to the algorithms within [rhoCOP](#).

The third L-comoment ratio matrix (L-skews and L-coskews) is

```

print(theoLC1cm$T3)
      [,1]      [,2]
[1,] 3.021969e-06 -2.829783e-05
[2,] -7.501135e-01 4.518901e-06

```

where the diagonal by definition should have zero zeros because the univariate L-skew of a uniform variable is zero. These values further support that default n is large enough. The secondary diagonal holds L-coskews. The copula has L-coskew of U wrt V of numerically near zero (symmetry) but measurable asymmetry of L-coskew of V wrt U of $\tau_3^{[21]} \approx -0.75$.

The fourth L-comoment ratio matrix (L-kurtosis and L-cokurtosis) is

```
print(theoLC1cm$T4)
      [,1]      [,2]
[1,] -2.623665e-06 -3.325177e-05
[2,] -2.162954e-04 -2.811630e-06
```

where the diagonal by definition should have zero zeros because the univariate L-kurtosis of a uniform variable is zero—it has no peakedness. These values further support that default n is large enough. The secondary diagonal holds L-cokurtosis and are near zero for this particular copula.

The fifth L-comoment ratio matrix (unnamed) is

```
print(theoLC1cm$T5)
      [,1]      [,2]
[1,] 1.813344e-06 -1.296025e-04
[2,] 1.246436e-01 1.475012e-06
```

where the diagonal by definition should have zero zeros because the univariate L-kurtosis of a uniform variable is zero—such a random variable has no asymmetry. These values further support that default n is large enough. The secondary diagonal holds the fifth L-comoment ratios. The copula has $\tau_5^{[12]} = 0$ of U wrt V of numerically near zero (symmetry) but measurable fifth-order L-comoment asymmetry of V wrt U of $\tau_5^{[21]} \approx 0.125$.

Comparison of Sample and Theoretical L-comoments—The previous section shows theoretical values computed as $\tau_3^{[21]} \approx -0.75$ and $\tau_5^{[21]} \approx 0.125$ for the two L-comoments substantially away from zero. As sample L-comoments these values are $\text{samLC\$T3}[2,1] = \hat{\tau}_3^{[21]} \approx -0.751$ and $\text{samLC\$T5}[2,1] = \hat{\tau}_5^{[21]} \approx 0.123$. **CONCLUSION:** *The sample L-comoment algorithms in the **lmomco** package are validated.*

Author(s)

W.H. Asquith

References

- Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.
- Brahimi, B., Chebana, F., and Necir, A., 2015, Copula representation of bivariate L-moments—A new estimation method for multiparameter two-dimensional copula models: *Statistics*, v. 49, no. 3, pp. 497–521.
- Nelsen, R.B., 2006, *An introduction to copulas*: New York, Springer, 269 p.
- Serfling, R., and Xiao, P., 2007, A contribution to multivariate L-moments—L-comoment matrices: *Journal of Multivariate Analysis*, v. 98, pp. 1765–1781.

See Also

[lcomCOP](#), [uvlmoms](#)


```
blomatrixCOPiqr(cop=NULL, para=NULL, as.sample=FALSE, as.blomCOPss=TRUE,
               ctype=c("weibull", "hazen", "1/n",
                       "bernstein", "checkerboard"), ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
as.sample	A logical controlling whether an optional R data.frame in para is used to compute the $\hat{\beta}_C^\circ$ -matrix at which point the ctype argument will be passed to multiple calls of EMPIRCop ;
as.blomCOPss	A logical to trigger blomCOPss for each of the (u, v) locations where for the blomCOPss calls $(\beta_C^\circ(\mathbf{u}, \mathbf{v}))$: $u \mapsto (u, u) \mapsto \mathbf{u}$ and $v \mapsto (v, v) \mapsto \mathbf{v}$;
ctype	Argument of the same as EMPIRCop ; and
...	Additional arguments to pass to the copula.

Value

The matrix for β_C° is returned depending on whether the decile or quartile version has been called.

Author(s)

W.H. Asquith

See Also

[blomCOP](#), [blomCOPss](#)

Examples

```
round(blomatrixCOPdec(cop=P), digits=8);      round(blomatrixCOPiqr(cop=P), digits=8)
#           U|V=0.10 U|V=0.50 U|V=0.90      #           U|V=0.25 U|V=0.50 U|V=0.75
# U|V=0.90           0           0           0      # U|V=0.75           0           0           0
# U|V=0.50           0           0           0      # U|V=0.50           0           0           0
# U|V=0.10           0           0           0      # U|V=0.25           0           0           0

round(blomatrixCOPdec(cop=PSP, as.blomCOPss=TRUE), digits=8)
#           U|V=0.10   U|V=0.50   U|V=0.90
# U|V=0.90 0.4736842  0.8181818  0.5153268
# U|V=0.50 0.8181818  0.3333333  0.6459330
# U|V=0.10 0.8901099  0.4736842  0.1708292

round(blomatrixCOPdec(cop=PSP, as.blomCOPss=FALSE), digits=8)
#           U|V=0.10   U|V=0.50   U|V=0.90
# U|V=0.90 0.09890110 0.81818182 4.2631579
# U|V=0.50 0.05263158 0.33333333 0.8181818
# U|V=0.10 0.01010101 0.05263158 0.0989011

## Not run:
set.seed(1)
```

```

td <- c(0.10, 0.50, 0.90)
UVn <- simCOP(n=5000, cop=glueCOP, col=8,
              para=list(glue=0.4, cop1 =PLcop,          cop2=PLcop,
                        para1=PLpar(rho=-0.5), para2=PLpar(rho=+0.5)))
points(td, rep(td[3], 3), cex=2, lwd=2, pch=3, col="red")
points(td, rep(td[2], 3), cex=2, lwd=2, pch=3, col="red")
points(td, rep(td[1], 3), cex=2, lwd=2, pch=3, col="red")

print(blomatrixCOPdec(as.sample=TRUE, para=UVn, ctype="weibull"))
#           U|V=0.10 U|V=0.50 U|V=0.90
# U|V=0.90 -0.08222222 -0.580 -0.190
# U|V=0.50  0.30800000  0.112  0.264
# U|V=0.10  0.84000000  0.620  0.262

BMdn <- blomatrixCOPdec(cop=glueCOP,
                       para=list(glue=0.4, cop1=PLcop,          cop2=PLcop,
                                 para1=PLpar(rho=-0.5), para2=PLpar(rho=+0.5)))
print(round(BMdn, digits=8))
#           U|V=0.10  U|V=0.50  U|V=0.90
# U|V=0.90 -0.08110464 -0.5569028 -0.2053772
# U|V=0.50  0.33449815  0.1202744  0.2424668
# U|V=0.10  0.75217766  0.6013719  0.2424668
## End(Not run)

## Not run:
set.seed(1); nsim <- 2000
para.pop <- list( cop1=GHCop,          cop2=PLcop,  alpha=0.359,
                  para1=c(4.003, 1.099), para2=0.882,  beta=0.292)
UVs <- simCOP(nsim, cop=composite2COP, para=para.pop)
mtext("GIVEN THIS SAMPLE")
Rho <- rhoCOP(as.sample=TRUE, para=UVs) # Spearman Rho
BMn <- blomatrixCOPdec(as.sample=TRUE, para=UVs, ctype="weibull")

parafn <- function(k) {
  c(exp(k[1])+1, exp(k[2]), exp(k[3]), pnorm(k[4]), pnorm(k[5]))
}
parafn_list <- function(k) {
  k <- parafn(k)
  list(cop1=GHCop, para1=c(k[1], k[2]), alpha=k[4],
       cop2=PLcop, para2=k[4],          beta=k[5])
}
BLOM_ofun <- function(para, statmat=NULL, parafn=NULL, rho=NA) {
  para <- parafn(para)
  new.para <- list(cop1=GHCop, para1=para[1:2], alpha=para[4],
                  cop2=PLcop, para2=para[3],  beta=para[5])
  bm <- blomatrixCOPdec(cop=composite2COP, para=new.para)
  err <- sum((statmat - bm)^2) + (rhoCOP(cop=composite2COP, para=new.para) - rho)^2
  #print(c(para, err))
  return(err)
}

run1 <- function(graphics=TRUE, nsim=0) {

```

```

par.init <- c(log(1), log(1), log(1), qnorm(0.5), qnorm(0.5))
rt      <- optim(par.init, BLOM_ofun, statmat=BMn, parafn=parafn, rho=Rho)
para    <- parafn(rt$par)
para.fit <- list( cop1=GHcop,      cop2=PLcop,  alpha=para[4],
                  para1=para[1:2], para2=para[3], beta=para[5])
uv <- simCOP(nsim, cop=composite2COP, para=para.fit, graphics=graphics, col=2)
rmse <- round(rmseCOP(uv[,1], uv[,2], ctype="weibull",
                     cop=composite2COP, para=para.fit), digits=8)
if(graphics) mtext(paste0("RMSE(run1)=", rmse))
return(list(rmse=rmse, para=para.fit))
}
system.time(RUN1 <- run1(nsim=nsim))
par.init <- c(log(RUN1$para$para1[1]), log(RUN1$para$para1[2]),
              log(RUN1$para$para2), qnorm(RUN1$para$alpha), qnorm(RUN1$para$beta))

RMSE_ofun <- function(para, parafn=NULL) {
  para <- parafn(para)
  new.para <- list(cop1=GHcop, para1=para[1:2], alpha=para[4],
                  cop2=PLcop, para2=para[3], beta=para[5])
  new.rmse <- rmseCOP(UVs[,1], UVs[,2], cop=composite2COP, para=new.para)
  #print(c(para, new.rmse))
  return(new.rmse)
}
run2 <- function(graphics=TRUE, nsim=0, par.init=NULL) {
  if(is.null(par.init)) {
    par.init <- c(log(1), log(1), log(1), qnorm(0.5), qnorm(0.5))
  }
  rt      <- optim(par.init, RMSE_ofun, parafn=parafn)
  para    <- parafn(rt$par)
  para.fit <- list( cop1=GHcop,      cop2=PLcop,  alpha=para[4],
                  para1=para[1:2], para2=para[3], beta=para[5])
  uv <- simCOP(nsim, cop=composite2COP, para=para.fit, graphics=graphics, col=4)
  rmse <- round(rmseCOP(uv[,1], uv[,2], ctype="weibull",
                     cop=composite2COP, para=para.fit), digits=8)
  if(graphics) mtext(paste0("RMSE(run2)=", rmse))
  return(list(rmse=rmse, para=para.fit))
}
system.time(RUN2.1 <- run2(nsim=nsim, par.init=par.init))
system.time(RUN2.2 <- run2(nsim=nsim, par.init=NULL ))

GIVN <- c(para.pop$alpha, para.pop$para1, para.pop$beta, para.pop$para2)
FIT1  <- c(RUN1$para$alpha, RUN1$para$para1, RUN1$para$beta, RUN1$para$para2)
FIT1  <- round(FIT1, digits=3)
FIT2.1 <- c(RUN2.1$para$alpha, RUN2.1$para$para1, RUN2.1$para$beta, RUN2.1$para$para2)
FIT2.1 <- round(FIT2.1, digits=3)
FIT2.2 <- c(RUN2.2$para$alpha, RUN2.2$para$para1, RUN2.2$para$beta, RUN2.2$para$para2)
FIT2.2 <- round(FIT2.2, digits=3)
nms    <- c("what", "alpha", "para1_1", "para1_2", "beta", "para2")
GIVN   <- c("given", GIVN); FIT2.1 <- c("by_Blom1", FIT2.1)
FIT1   <- c("by_RMSE", FIT1); FIT2.2 <- c("by_Blom2", FIT2.2)
names(GIVN) <- nms; names(FIT1) <- nms
names(FIT2.1) <- nms; names(FIT2.2) <- nms
RESL <- cbind(data.frame(GIVN), data.frame(FIT1), data.frame(FIT2.1), data.frame(FIT2.2))

```

```
print(RESL) #
## End(Not run)
```

blomCOP

The Blomqvist Beta of a Copula

Description

Compute the *Blomqvist Beta* $\beta_{\mathbf{C}}$ of a copula (Nelsen, 2006, p. 182), which is defined at the middle or center of $\mathcal{I}^2$ as

$$\beta_{\mathbf{C}} = 4 \times \mathbf{C}\left(\frac{1}{2}, \frac{1}{2}\right) - 1,$$

where the $u = v = 1/2$ and thus shows that $\beta_{\mathbf{C}}$ is based on the *median joint probability*. The Blomqvist Beta is also called the *medial correlation coefficient*. Nelsen also reports that “although, the Blomqvist Beta depends only on the copula only through its value at the center of $\mathcal{I}^2$, but that $[\beta_{\mathbf{C}}]$ nevertheless often provides an accurate approximation to both *Spearman Rho* [rhoCOP](#) and *Kendall Tau* [tauCOP](#).” Kendall Tau $\tau_{\mathbf{C}}$, *Gini Gamma* $\gamma_{\mathbf{C}}$, and Spearman Rho $\rho_{\mathbf{C}}$ in relation to $\beta_{\mathbf{C}}$ satisfy the following inequalities (Nelsen, 2006, exer. 5.17, p. 185):

$$\begin{aligned} \frac{1}{4}(1 + \beta_{\mathbf{C}})^2 - 1 &\leq \tau_{\mathbf{C}} \leq 1 - \frac{1}{4}(1 - \beta_{\mathbf{C}})^2, \\ \frac{3}{16}(1 + \beta_{\mathbf{C}})^3 - 1 &\leq \rho_{\mathbf{C}} \leq 1 - \frac{3}{16}(1 - \beta_{\mathbf{C}})^3, \text{ and} \\ \frac{3}{8}(1 + \beta_{\mathbf{C}})^2 - 1 &\leq \tau_{\mathbf{C}} \leq 1 - \frac{3}{8}(1 - \beta_{\mathbf{C}})^2. \end{aligned}$$

A curious aside (Joe, 2014, p. 164) about the *Gaussian copula* is that *Blomqvist Beta* is equal to *Kendall Tau* ([tauCOP](#)): $\beta_{\mathbf{C}} = \tau_{\mathbf{C}}$ (see **Note** in [med.regressCOP](#) for a demonstration). Finally, a version of Blomqvist Beta defined outside the median is provided by [blomCOPss](#).

Usage

```
blomCOP(cop=NULL, para=NULL, as.sample=FALSE,
        ctype=c("joe", "weibull", "hazen", "1/n",
                 "bernstein", "checkerboard"), ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
as.sample	A logical controlling whether an optional R data.frame in para is used to compute the $\hat{\beta}_{\mathbf{C}}$ (see Note);
ctype	Argument of the same as EMPIRCOP with the exception of the “joe” specific to the documentation here. The other choices trigger and are given over to the empirical copula; and
...	Additional arguments to pass to the copula or down to EMPIRCOP if a sample version had been requested.

Value

The value for β_C or sample $\hat{\beta}_n$ is returned.

Note

The sample $\hat{\beta}_n$ is most efficiently computed (Joe, 2014, p. 57) by

$$\hat{\beta}_n = \frac{2}{n} \sum_{i=1}^n \mathbf{1} \left([r_{i1} - (1+n)/2] \times [r_{i2} - (1+n)/2] \geq 0 \right) - 1,$$

where r_{i1}, r_{i2} are the ranks of the data for $i = 1, \dots, n$, and $\mathbf{1}(\cdot)$ is an *indicator function* scoring 1 if condition is true otherwise zero. However, the Joe sample estimator is not fully consistent (or vice versa) with the various versions of the empirical copula, C_n , ([EMPIRCop](#)) (see the last example in **Examples**). Also, the nature of even and odd sample sizes controls how the median is computed and the issue of samples lying on the median lines in U and V (Genest *et al.*, 2013). The argument `cotype` supports triggers to the C_n in lieu of the Joe sample estimator shown in this documentation.

Author(s)

W.H. Asquith

References

Genest, C., Carabarin-Aguirre, A., and Harvey, F., 2013, Copula parameter estimation using Blomqvist's beta: Journal de la Société Française de Statistique, v. 154, no. 1, pp. 5–24.

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[blomCOPss](#), [blomatrixCOPdec](#), [blomatrixCOPiqr](#), [footCOP](#), [giniCOP](#), [hoefCOP](#), [rhoCOP](#), [tauCOP](#), [wolfCOP](#), [joeskewCOP](#), [uvmoms](#)

Examples

```
blomCOP(cop=PSP) # 1/3 precisely

## Not run:
# Nelsen (2006, exer. 5.17, p. 185) : All if(...) are TRUE
B <- blomCOP(cop=N4212cop, para=2.2); Bp1 <- 1 + B; Bm1 <- 1 - B
G <- giniCOP(cop=N4212cop, para=2.2); a <- 1/4; b <- 3/16; c <- 3/8
R <- rhoCOP(cop=N4212cop, para=2.2)
K <- tauCOP(cop=N4212cop, para=2.2, brute=TRUE) # numerical issues without brute
if( a*Bp1^2 - 1 <= K & K <= 1 - a*Bm1^2 ) print("TRUE") #
if( b*Bp1^3 - 1 <= R & R <= 1 - b*Bm1^3 ) print("TRUE") #
if( c*Bp1^2 - 1 <= G & G <= 1 - c*Bm1^2 ) print("TRUE") #
## End(Not run)

## Not run:
# A demonstration of a special feature of blomCOP for sample data.
```

```

# Joe (2014, p. 60; table 60) has 0.749 for GHcop(tau=0.5); n*var(hatB) as n-->infinity
set.seed(1)
theta <- GHcop(tau=0.5)$para; B <- blomCOP(cop=GHcop, para=theta); n <- 1000
H <- sapply(1:1000, function(i) { # Let us test that with pretty large sample size:
    blomCOP(para=rCOP(n=n, cop=GHcop, para=theta), as.sample=TRUE) })
print(n*var(B-H)) # For 1,000 sims of size n : 0.789, nearly matches Joe's result
## End(Not run)

## Not run:
# Joe (2014, p. 57) says that sqrt(n)(B-HatBeta) is Norm(0, 1 - B^2)
set.seed(1)
n <- 10000; B <- blomCOP(cop=PSP) # Beta = 1/3
H <- sapply(1:100, function(i) { message(i,"-", appendLF=FALSE)
    blomCOP(para=rCOP(n=n, cop=PSP), as.sample=TRUE) })
lmomco::parnor(lmomco::lmoms(sqrt(n)*(H-B))) # mu = -0.038; sigma = 0.970
# Joe (2014) : sqrt(1-B^2) == standard deviation (sigma) : (1-(1/3)^2) approx 0.973
## End(Not run)

## Not run:
nn <- 200; set.seed(1)
UV <- simCOP(n=nn+1, cop=PSP, graphics=FALSE)
for(n in nn:(nn+1)) {
  if(as.logical(n %% 2)) { # in source \ percent \ percent for latex
    message("Blomquist Betas for an odd sample size n=", n)
    uv <- UV
  } else {
    message("Blomquist Betas for an even sample size n=", n)
    uv <- UV[-(nn+1), ] # remove the last and 'odd' indexed value to make even
  }
  message(c(" Joe2014: ", blomCOP(as.sample=TRUE, para=uv, ctype="joe"      )))
  message(c(" Weibull: ", blomCOP(as.sample=TRUE, para=uv, ctype="weibull"  )))
  message(c(" Hazen:   ", blomCOP(as.sample=TRUE, para=uv, ctype="hazen"    )))
  message(c(" 1/n:    ", blomCOP(as.sample=TRUE, para=uv, ctype="1/n"      )))
  message(c(" Bernstn: ", blomCOP(as.sample=TRUE, para=uv, ctype="bernstein" )))
  message(c(" ChckBrd: ", blomCOP(as.sample=TRUE, para=uv, ctype="checkerboard"))))
}
# Blomquist Betas for an even sample size n=200
# Joe2014: 0.32
# Weibull: 0.32
# Hazen:   0.32
# 1/n:     0.32
# Bernstn: 0.323671819423416
# ChckBrd: 0.32
# Blomquist Betas for an odd sample size n=201
# Joe2014: 0.323383084577114
# Weibull: 0.333333333333333
# Hazen:   0.333333333333333
# 1/n:     0.293532338308458
# Bernstn: 0.327747577290946
# ChckBrd: 0.313432835820896 #
## End(Not run)

```

Description

Compute the *Blomqvist (Schmid–Schmidt) Betas* $\beta_C^\diamond$ (Schmid and Schmidt, 2007) defined for arbitrary dimension d of a copula $\mathbf{C}(u_1, \dots, u_d; \Theta)$ (COP) for parameters Θ . The copula survival function is $\overline{\mathbf{C}}(u_1, \dots, u_d; \Theta)$ (surfuncCOP). The Beta, though the **copBasic** package is built around bivariate copula only, is defined as

$$\beta_C^\diamond = h_d(\mathbf{u}, \mathbf{v}) [(\mathbf{C}(\mathbf{u}) + \overline{\mathbf{C}}(\mathbf{v})) - g_d(\mathbf{u}, \mathbf{v})],$$

where h_d and g_d are norming constants defined below. The superscript $\diamond$ (diamond) is chosen for **copBasic** because of the alliteration to “dimension.” The bold face font for $\mathbf{u}$ and $\mathbf{v}$ shows these arguments as vectors of length d reflecting “cutting points” on nonexceedance probabilities in each of the dimensions. The $\mathbf{u}$ functions as the arguments (u, v) pair used in copula of this package and represents the first cutting point for a $\Pr[U \leq u, V \leq v] = \mathbf{C}(u, v)$, and $\mathbf{v}$ functions as the arguments u, v pair for this package and represents the second cutting point for a $\Pr[U > u, V > v] = 1 - u - v + \mathbf{C}(u, v) = \overline{\mathbf{C}}(u, v)$. This notation of vectored (bold face) and nonvectored “ u ” and “ v ” is a little obtuse but as the properties of $\beta_C^\diamond$ are summarized clarity for the reader is anticipated. In short, the $\mathbf{u}$ will reference the coordinate pairs in the lower right quadrant and the $\mathbf{v}$ will reference the coordinate pairs in the upper right quadrant.

The norming constant h_d is defined as

$$h_d(\mathbf{u}, \mathbf{v}) = \frac{1}{(\min(u_1, \dots, u_d) + \min(1 - v_1, \dots, 1 - v_d) - g_d(\mathbf{u}, \mathbf{v}))},$$

and g_d is defined as

$$g_d(\mathbf{u}, \mathbf{v}) = \prod_{i=1}^d u_i + \prod_{i=1}^d (1 - v_i),$$

where the cutting points $\mathbf{u}$ and $\mathbf{v}$ are in a domain $D : \{(\mathbf{u}, \mathbf{v})\} \in [0, 1]^{2d}$ given $\mathbf{u} \leq \mathbf{v}$ and $\mathbf{u} > 0$ or $\mathbf{v} < 1$. The reader must careful remember that these $\mathbf{u}$ and $\mathbf{v}$ are vectors of probabilities.

The norming constants provide for $-1 \leq \beta_C^\diamond \leq +1$. Using the function argument defaults for $d = 2$ dimensions $\mathbf{u} = (1, 1)/2$ for uu and $\mathbf{v} = (1, 1)/2$ for vv , results in (1) $\beta_C^\diamond = 1$ if $\mathbf{C} = \mathbf{M}$ *comonotonicity copula* (M) (blomCOPss(cop=M) == 1), (2) $\beta_C^\diamond = 0$ if $\mathbf{C} = \mathbf{P}$ *independence copula* (P) (blomCOPss(cop=P) == 0), and (3) if $\mathbf{C} = \mathbf{W}$ *countermonotonicity copula* (W) $\beta_C^\diamond = 1$ (blomCOPss(cop=W) == -1).

Schmid and Schmidt (2007) list three important cases extending the **M** and **P** examples. First, $\beta_C^\diamond(1/2, 1/2) = \beta_C(1/2, 1/2)$, which is *Blomqvist Beta* ($\beta_C(1/2, 1/2)$) (blomCOP) and measures overall dependence.

Second, $\beta_C^\diamond(\mathbf{u}, \mathbf{v})$ with $\mathbf{u} < 1/2 < \mathbf{v}$, which measures dependence in the tail regions. (Note, the author of **copBasic** thinks “regions” as a plural is need in the previous sentence; Schmid and Schmidt (2007) use the singular “region.” This is potentially important as seemingly simultaneous tail dependency in the lower and upper perspectives would be provided. More discussion is provided in **Examples**.)

Third and presumably very important in practical applications, $\lim_{p \downarrow 0} \beta_{\mathbf{C}}^{\circ}(\mathbf{p}, \mathbf{1}) = \lambda_{\beta_{\mathbf{C}}^{\circ}}^L$ for $\mathbf{p} = \mathbf{u} = (p, \dots, p)$ measures lower-tail dependence. This measure is equal to the *lower-tail dependence parameter* $\lambda_{\beta_{\mathbf{C}}^{\circ}}^L = \lambda_{\beta_{\mathbf{C}}^{\circ}}^L$ without some of the computational nuances required as $\lambda_{\mathbf{C}}^L$ is defined at [taildepCOP](#).

Schmid and Schmidt (2007) do not list how the *upper-tail dependence parameter* $\lambda_{\mathbf{C}}^U$ could be computed in terms of $\beta_{\mathbf{C}}^{\circ}$. The expression for study of the upper-tail dependency is $\lambda_{\beta_{\mathbf{C}}^{\circ}}^U = \beta_{\mathbf{C}}^{\circ}(\mathbf{0}, \mathbf{p})$ for $\mathbf{p} = \mathbf{v} = (p, \dots, p)$ as $p \rightarrow 0^+$, and $\lambda_{\mathbf{C}}^U = \lambda_{\beta_{\mathbf{C}}^{\circ}}^U$ without some of the computational nuances required as $\lambda_{\mathbf{C}}^U$ is defined at [taildepCOP](#). These tail dependencies are computed and compared in the **Examples** and confirmation of this function being used to estimate both tail-dependency parameters is confirmed.

Usage

```
blomCOPss(cop=NULL, para=NULL, uu=rep(0.5, 2), vv=rep(0.5, 2), trap.nan=TRUE,
          as.sample=FALSE, ctype=c("weibull", "hazen", "1/n",
                                   "bernstein", "checkerboard"), ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
uu	The vector for $\mathbf{u}$ and the defaults with $\mathbf{vv}$ as such for same operation as blomCOP ($\beta_{\mathbf{C}}^{\circ}(\mathbf{1}/2, \mathbf{1}/2)$);
vv	The vector for $\mathbf{v}$ and the defaults with $\mathbf{uu}$ as such for same operation as blomCOP ($\beta_{\mathbf{C}}^{\circ}(\mathbf{1}/2, \mathbf{1}/2)$);
trap.nan	A logical to trigger 0 if (0, 0) is NaN or if (1, 1) is NaN. This feature is present on a package-specific purpose because the PSP copula deliberately retains edge NaN as a stress case;
as.sample	A logical controlling whether an optional R data.frame in para is used to compute the $\hat{\beta}_{\mathbf{C}}^{\circ}$ at which point the ctype argument will be passed to EMPIRCop ;
ctype	Argument of the same as EMPIRCop ; and
...	Additional arguments to pass to the copula.

Value

The $\beta_{\mathbf{C}}^{\circ}$ is returned.

Note

Sample estimation of the $\beta_{\mathbf{C}}^{\circ}$ is possible. The `as.sample` triggers internally a call to the *empirical copula* ($\mathbf{C}_n$) ([EMPIRCop](#)) for the `ctype` for the copula and its survival function form. Expansive more details are provided by [taildepCOP](#) (section **Note::DEMONSTRATION (Tail Dependence)**). A comparison of the $\hat{\lambda}_{\beta_{\mathbf{C}}^{\circ}}^U$ and $\hat{\lambda}_{\beta_{\mathbf{C}}^{\circ}}^L$ is made.

Author(s)

W.H. Asquith

References

Schmid, Friedrich, and Schmidt, Rafael, 2007, Nonparametric inference on multivariate versions of Blomqvist's beta and related measures of tail dependence: *Metrika*, v. 66, pp. 323–354, [doi:10.1007/s0018400601143](https://doi.org/10.1007/s0018400601143).

See Also

[blomCOP](#), [blomatrixCOP](#), [taildepCOP](#)

Examples

```
blomCOP( cop=PSP) # [1] 0.3333333
blomCOPss(cop=PSP) # [1] 0.3333333

## Not run:
# The calls below for blomCOPss() are technically the same for sample versions.
UV <- simCOP(1000, cop=PSP, graphics=FALSE) # HatBeta(0.1,0.9) = 0.277___
blomCOPss(para=UV, cop=EMPIRCop, uu=c(0.1,0.1), vv=c(0.90,0.90))
blomCOPss(para=UV, as.sample=TRUE, uu=c(0.1,0.1), vv=c(0.90,0.90)) #
## End(Not run)

## Not run:
set.seed(1)
para <- c(3, 6) # define parameters of two-parameter GHcop
UV <- simCOP(1000, cop=GHcop, para=para) # simulate to show general structure

# compute the tail dependencies from having into the limits
taildepCOP(cop=GHcop, para=para, plot=TRUE)
# lower tail dependency = 0.96222
# upper tail dependency = 0.74008
# The two parameters influence how strongly the tail dependencies are.

# Schmid and Schmidt (2007, eq. 24) define the lower-tail dependency in terms of
# the Beta and  $p \rightarrow 0$  Beta( $c(p,p)$ ,  $c(1,1)$ ). Lets compute these and produce content
# suitable to show on the tail-dependency plot that the assertion for the lower
# dependency by Beta() is correct, which it is and then extend to the upper-tail
# dependency parameter that the authors seem to not have defined.
usr <- par()$usr[1:2] # grab horizontal edges of the plot, and set up the
uuL0 <- rep(pnorm(usr[1]), 2) # the uu for the lower tail and the vv for the upper
vvUP <- rep(pnorm(usr[2]), 2) # tail and then plot both with overplotting symbols
# lower-tail estimate and see how it plots along the value from taildepCOP()
SchmidsL <- blomCOPss(cop=GHcop, para=para, uu=uuL0, vv=c(1,1))
points(usr[1], SchmidsL, col="darkgreen", cex=2, pch=1, lwd=2)
points(usr[1], SchmidsL, col="darkgreen", cex=2, pch=3, lwd=2)
points(usr[1], SchmidsL, col="darkgreen", cex=2, pch=4, lwd=2)
# upper-tail estimate and see how it plots along the value from taildepCOP()
SchmidsU <- blomCOPss(cop=GHcop, para=para, uu=c(0,0), vv=vvUP)
points(usr[2], SchmidsU, col="darkgreen", cex=2, pch=1, lwd=2)
points(usr[2], SchmidsU, col="darkgreen", cex=2, pch=3, lwd=2)
points(usr[2], SchmidsU, col="darkgreen", cex=2, pch=4, lwd=2)
# SchmidsL lower tail dependency = 0.962224
# SchmidsU upper tail dependency = 0.740079
```

```

# The author has an expectation that the SchmidSL and SchmidSU values are
# more reliable than those stemming from taildepCOP() because of the limiting
# behavior (or its implementation therein) compared to direct computation by
# blomCOPss().

# Now for sake of curiosity, let us see how the trajectory of the Blomqvist
# (Schmid--Schmidt) Betas at arriving at the tail dependencies as p-->0|1.
# It is very informative that the trajectories of blomCOPss() and taildepCOP()
# as each hones towards the two dependency parameters are different and this
# highlights the fact that the computational underpinnings are different.
psl <- pnorm(seq(0, usr[1], by=-diff(range(c(0, usr[1])))) / 1000))
lines(qnorm(psl), sapply(psl, function(p) {
  blomCOPss(cop=GHcop, para=para, uu=rep(p, 2), vv=c(1,1)) } ),
  col="darkgreen", lty=2, lwd=2)
psu <- pnorm(seq(0, usr[2], by= diff(range(c(0, usr[2])))) / 1000))
lines(qnorm(psu), sapply(psu, function(p) {
  blomCOPss(cop=GHcop, para=para, uu=c(0,0), vv=rep(p, 2)) } ),
  col="darkgreen", lty=2, lwd=2) #
## End(Not run)

```

breveCOP

Add Asymmetry to a Copula

Description

Adding *permutation asymmetry* (Chang and Joe, 2020, p. 1596) ([isCOP.permsym](#)) is simple for a bivariate copula family. Let $\mathbf{C}$ be a copula with respective vectors of parameters $\Theta_{\mathbf{C}}$, then the permutation asymmetry is added through an asymmetry parameter $\beta \in (-1, +1)$ by

$$\check{\mathbf{C}}_{\beta;\Theta}(u, v) = v^{-\beta} \cdot \mathbf{C}(u, v^{(1+\beta)}; \Theta), \text{ and}$$

for $0 \leq \beta \leq +1$ by

$$\check{\mathbf{C}}_{\beta;\Theta}(u, v) = u^{+\beta} \cdot \mathbf{C}(u^{(1-\beta)}, v; \Theta).$$

The parameter β clashes in name and symbology with a parameter used by functions [composite1COP](#), [composite2COP](#), and [composite3COP](#). As a result, support for alternative naming is provided for compatibility. Lastly, an example under [W](#) demonstrates the span of the *Kendall function* ([kfuncCOP](#)) based on [M](#) and [W](#) having permutation asymmetry inserted by [breveCOP\(\)](#). The demonstration shows that the Kendall function is a distribution unto itself with its own nonexceedance probabilities.

Usage

```
breveCOP(u,v, para, breve=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A special parameter list (see Note);
<code>breve</code>	An alternative way from <code>para</code> to set the β for this function; and
<code>...</code>	Additional arguments to pass to the copula.

Value

Value(s) for the copula are returned.

Note

The following descriptions list in detail the structure and content of the `para` argument where `cop1` and `cop` and `para1` and `para` are respectively synonymous to have some structural similarity to the various copula constructors (compositors) of the **copBasic** package:

`beta` — The β asymmetry parameter;

`breve` — The β asymmetry parameter and presence of `breve` will not cause the non-use of `beta`; this feature is present so that `beta` remains accessible to the compositors that use `beta` (see **Examples**);

`cop` — Function of the copula $\mathbf{C}$;

`cop1` — Alternative naming of the function of the coupla $\mathbf{C}$;

`para` — Vector of parameters $\Theta_{\mathbf{C}}$ for $\mathbf{C}$; and

`para1` — Alternative naming of the vector of parameters $\Theta_{\mathbf{C}}$ for $\mathbf{C}$.

The function silently restricts the β to its interval as defined, but parameter transform might be useful in some numerical optimization schemes. The following recipes might be useful for transform from a parameter in numerical optimization to the asymmetry parameter:

```
# transform into space for optimization
BREVEtfunc <- function(p) { return( qnorm((p[1] + 1) / 2) ) } # [-Inf,+Inf]
# re-transform back into space for the copula
BREVErfunc <- function(p) { return(2*pnorm(p[1]) - 1) } # [-1 ,+1 ]
```

Author(s)

W.H. Asquith

References

Chang, B., and Joe, H., 2020, Copula diagnostics for asymmetries and conditional dependence: Journal of Applied Statistics, v. 47, no. 9, pp. 1587–1615, doi:10.1080/02664763.2019.1685080.

See Also

[COP](#), [convex2COP](#), [convexCOP](#), [composite1COP](#), [composite2COP](#), [composite3COP](#), [FRECHETcop](#), [glueCOP](#)

Examples

```

para <- list(breve=0.24, cop1=FRECHETcop, para1=c(0.4, 0.56))
breveCOP(0.87, 0.35, para=para) # 0.282743

betas <- seq(-1,1, by=0.01)
bloms <- sapply(betas, function(b) {
  breveCOP(0.15, 0.25, para=list(cop=GLPMcop, para=c(2, 2), beta=b))
})
plot(betas, bloms, type="l", main="GLPMcop(u,v; 2,2) by breveCOP(beta)")

## Not run:
# Notice the argument cop and para name adjustments to show that
# translation exists inside the function to have use flexibility.
para <- list(breve=+0.44, cop1=FRECHETcop, para1=c(0.2, 0.56))
UV <- simCOP(1000, cop=breveCOP, para=para)
para <- list(breve=-0.44, cop= FRECHETcop, para= c(0.2, 0.56))
UV <- simCOP(1000, cop=breveCOP, para=para) #
## End(Not run)

## Not run:
# Testing on a comprehensive copula (Plackett)
betas <- rhos <- thetas <- brhos <- NULL
for(beta in seq(-1, 1, by=0.1 )) {
  for(rho in seq(-1, 1, by=0.01)) {
    theta <- PLACKETTpar(rho=rho, byrho=TRUE)
    thetas <- c(thetas, theta)
    para <- list( cop=PLcop, para=theta, beta=beta)
    brho <- rhoCOP(cop=breveCOP, para=para)
    betas <- c(betas, beta); rhos <- c(rhos, rho)
    brhos <- c(brhos, brho)
  }
}
df <- data.frame(beta=betas, theta=thetas, rho=rhos, brho=brhos)
plot(df$theta, df$brho, log="x", pch=16, cex=0.9, col="seagreen",
      xlab="Plackett parameter", ylab="Spearman Rho")
lines(df$theta[df$beta == 0], df$brho[df$beta == 0], col="red", lwd=2)
# Red line is the Plackett in its permutation symmetric definition. #
## End(Not run)

## Not run:
# Here is an example for a test using mleCOP() to estimate a 5-parameter asymmetric
# copula model to "some data" on transition from yesterday to today data for a very
# large daily time series. The purpose of example here is to demonstrate interfacing
# to the breveCOP() for it to add asymmetry to composition of two copula.
myASYMCOP <- function(u,v, para, ...) {
  subpara <- list(alpha=para$alpha, beta=para$beta, cop1=GHcop, para1=para$para1,
                  cop2=PLcop, para2=para$para2)

  breveCOP(u,v, cop=convex2COP, para=subpara)
}
para <- list(alpha=+0.16934027, cop1=GHcop, para1=c(1.11144148, 10.32292673),
            beta=-0.01923808, cop2=PLcop, para2=3721.82966727)
UV <- simCOP(30000, cop=myASYMCOP, para=para, pch=16, col=grey(0, 0.1))

```

```

    abline(0,1, lwd=3, col="red") #
## End(Not run)

## Not run:
# Here is a demonstration of the permutations of the passing of the
# asymmetry parameter into the function and then by
UV <- simCOP(1E3, cop=breveCOP, para=list(cop=HRcop, para=5), breve=+0.5)
UV <- simCOP(1E3, cop=breveCOP, para=list(cop=HRcop, para=5), breve=-0.5)
UV <- simCOP(1E3, cop=breveCOP, para=list(cop=HRcop, para=5, beta =+0.5))
UV <- simCOP(1E3, cop=breveCOP, para=list(cop=HRcop, para=5, breve=+0.5))
UV <- simCOP(1E3, cop=breveCOP, para=list(cop=HRcop, para=5, beta=-0.4, breve=+0.5))

para <- list(cop1=HRcop, para1=6, cop2=PSP, para2=NULL, alpha=1, beta=0.7)
myCOP <- function(u,v, para, ...) breveCOP(u,v, cop=composite2COP, para=para)
para$breve <- "here I am"
UV <- simCOP(1E3, cop=composite2COP, para=para, seed=1) # breve is not used
para$breve <- -0.16
UV <- simCOP(1E3, cop=myCOP, para=para, seed=1)
para$breve <- +0.16
UV <- simCOP(1E3, cop=myCOP, para=para, seed=1) #
## End(Not run)

```

CIRCcop

Copula of Circular Uniform Distribution

Description

The *Circular copula* of the coordinates (x, y) of a point chosen at random on the *unit circle* (Nelsen, 2006, p. 56) is given by

$$\mathbf{C}_{\text{CIRC}}(u, v) = \mathbf{M}(u, v) \text{ for } |u - v| > 1/2,$$

$$\mathbf{C}_{\text{CIRC}}(u, v) = \mathbf{W}(u, v) \text{ for } |u + v - 1| > 1/2, \text{ and}$$

$$\mathbf{C}_{\text{CIRC}}(u, v) = \frac{u + v}{2} - \frac{1}{4} \text{ otherwise .}$$

The coordinates of the unit circle are given by

$$\mathbf{CIRC}(x, y) = \left(\frac{\cos(\pi(u - 1)) + 1}{2}, \frac{\cos(\pi(v - 1)) + 1}{2} \right).$$

Usage

```
CIRCcop(u, v, para=NULL, as.circ=FALSE, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	Optional parameter list argument that can contain the logical <code>as.circ</code> instead;
<code>as.circ</code>	A logical, if true, to trigger the transformation $u = 1 - \text{acos}(2x - 1)/\pi$ and $v = 1 - \text{acos}(2y - 1)/\pi$ to convert (X, Y) coordinates of a uniform unit circle to the (U, V) in nonexceedance probability; and
<code>...</code>	Additional arguments to pass, if ever needed.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

Examples

```

CIRCcop(0.5, 0.5) # 0.25 quarterway along the diagonal upward to right
CIRCcop(0.5, 1 ) # 0.50 halfway across in horizontal direction
CIRCcop(1 , 0.5) # 0.50 halfway across in vertical direction

## Not run:
nsim <- 2000
opts <- par(xpd=NA, las=1, lend=2, no.readonly=TRUE)
rtheta <- runif(nsim, min=0, max=2*pi) # polar coordinate simulation
XY <- data.frame(X=cos(rtheta)/2 + 1/2, Y=sin(rtheta)/2 + 1/2)
plot(XY, lwd=0.8, col="lightgreen", xaxs="i", yaxs="i",
      xlab="X OF UNIT CIRCLE OR NONEXCEEDANCE PROBABILITY U",
      ylab="Y OF UNIT CIRCLE OR NONEXCEEDANCE PROBABILITY V")
UV <- simCOP(nsim, cop=CIRCcop, lwd=0.8, col="salmon3", ploton=FALSE)
theta <- 3/4*pi+0.1 # select a point on the upper left of the circle
x <- cos(theta)/2 + 1/2; y <- sin(theta)/2 + 1/2 # coordinates
H <- CIRCcop(x, y, as.circ=TRUE) # 0.218169 # Pr[X <= x & Y <= y]
points(x, y, pch=16, col="forestgreen", cex=2)
segments(0, y, x, y, lty=2, lwd=2, col="forestgreen")
segments(x, 0, x, y, lty=2, lwd=2, col="forestgreen")
Hemp1 <- sum(XY$X <= x & XY$Y <= y) / nrow(XY) # about 0.22 as expected
u <- 1-acos(2*x-1)/pi; v <- 1-acos(2*y-1)/pi
segments(0, v, u, v, lty=2, lwd=2, col="salmon3")
segments(u, 0, u, v, lty=2, lwd=2, col="salmon3")
points(u, v, pch=16, cex=2, col="salmon3")
arrows(x, y, u, v, code=2, lwd=2, angle=15) # arrow points from (X,Y) coordinate
# specified by angle theta in radians on the unit circle to the corresponding
# coordinate in (U,V) domain of uniform circular distribution copula

```

[illegible]

```

ylab=paste0("Y OF CIRCULAR UNIFORM DISTRIBUTION OR\n",
            "NONEXCEEDANCE PROBABILITY OF V"))
JK <- data.frame(U=1 - acos(2*XY$X - 1)/pi, V=1 - acos(2*XY$Y - 1)/pi)
segments(x0=UV$U, y0=UV$V, x1=XY$X, y1=XY$Y, col="lightgreen", lwd=0.8)
points(XY, lwd=0.8, col="darkgreen")
points(JK, pch=16, col="salmon3", cex=0.5)

t <- seq(0.001, 0.999, by=0.001)
t <- diagCOPatf(t, cop=composite2COP, para=para)
AB <- data.frame(X=(cos(pi*(t-1))+1)/2, Y=(cos(pi*(t-1))+1)/2)
lines(AB, lwd=4, col="seagreen") #
## End(Not run)

```

CLcop

The Clayton Copula

Description

The *Clayton copula* (Joe, 2014, p. 168) is

$$\mathbf{C}_{\Theta}(u, v) = \mathbf{CL}(u, v) = \max[(u^{-\Theta} + v^{-\Theta} - 1; 0)]^{-1/\Theta},$$

where $\Theta \in [-1, \infty)$, $\Theta \neq 0$. The copula, as $\Theta \rightarrow -1^+$ limits, to the *countermonotonicity copula* ($\mathbf{W}(u, v)$; [W](#)), as $\Theta \rightarrow 0$ limits to the *independence copula* ($\mathbf{\Pi}(u, v)$; [P](#)), and as $\Theta \rightarrow \infty$, limits to the *comonotonicity copula* ($\mathbf{M}(u, v)$; [M](#)). The parameter Θ is readily computed from a *Kendall Tau* ([tauCOP](#)) by $\tau_{\mathbf{C}} = \Theta/(\Theta + 2)$.

Usage

```
CLcop(u, v, para=NULL, tau=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula;
<code>tau</code>	Optional Kendall Tau; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned. Otherwise if `tau` is given, then the Θ is computed and a list having

<code>para</code>	The parameter Θ , and
<code>tau</code>	Kendall Tau.

and if `para=NULL` and `tau=NULL`, then the values within `u` and `v` are used to compute Kendall Tau and then compute the parameter, and these are returned in the aforementioned list.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also[M](#), [P](#), [W](#)**Examples**

```
# Lower tail dependency of Theta = pi --> 2^(-1/pi) = 0.8020089 (Joe, 2014, p. 168)
taildepCOP(cop=CLcop, para=pi)$lambdaL # 0.80201
```

coCOP

*The Co-Copula Function***Description**

Compute the *co-copula (function)* from a copula (Nelsen, 2006, pp. 33–34), which is defined as

$$\Pr[U > u \text{ or } V > v] = \mathbf{C}^*(u', v') = 1 - \mathbf{C}(u', v'),$$

where $\mathbf{C}^*(u', v')$ is the co-copula and u' and v' are exceedance probabilities and are equivalent to $1 - u$ and $1 - v$ respectively. The co-copula is the expression for the probability that either $U > u$ or $V > v$ when the arguments to $\mathbf{C}^*(u', v')$ are exceedance probabilities, which is unlike the *dual of a copula (function)* (see [duCOP](#)) that provides $\Pr[U \leq u \text{ or } V \leq v]$.

The co-copula is a function and not in itself a copula. Some rules of copulas mean that $\mathbf{C}(u, v) + \mathbf{C}^*(u', v') \equiv 1$ or in **copBasic** syntax that the functions $\text{COP}(u, v) + \text{coCOP}(u, v)$ equal unity if the exceedance argument to coCOP is set to FALSE.

The function coCOP gives “risk” against failure if failure is defined as either hazard source U or V occurring by themselves or if both occurred at the same time. Expressing this in terms of an annual probability of occurrence (q), one has

$$q = 1 - \Pr[U > u \text{ or } V > v] = \mathbf{C}^*(u', v') \text{ or}$$

in R code `q <- coCOP(u, v, exceedance=FALSE, ...)`. So, in yet other words and as a mnemonic: *A co-copula is the probability of exceedance if the hazard sources **collaborate** or **cooperate** to cause failure.* Also, q can be computed by `q <- coCOP(1 - u, 1 - v, exceedance=TRUE, ...)`.

Usage

```
coCOP(u, v, cop=NULL, para=NULL, exceedance=TRUE, ...)
```

Arguments

<code>u</code>	Exceedance probability ($u' = 1 - u$) in the X direction;
<code>v</code>	Exceedance probability ($v' = 1 - v$) in the Y direction;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>exceedance</code>	A logical controlling the probability direction. Are <code>u</code> and <code>v</code> values given really u' and v' , respectively? If FALSE, then the complements of the two are made internally and the nonexceedances can thus be passed; and
<code>...</code>	Additional arguments to pass to the copula.

Value

The value(s) for the co-copula are returned.

Note

The author (Asquith) finds the use of exceedance probabilities delicate in regards to Nelsen's notation. The `coCOP` function and `surCOP` have the exceedance argument to serve as a reminder that the co-copula as defined in the literature uses *exceedance probabilities* as its arguments, although the arguments as code `u` and `v` do not mimic the overline nomenclature ($\overline{\cdot}$) of the exceedance (survival) probabilities.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

`COP`, `surCOP`, `duCOP`

Examples

```
u <- 1 - runif(1); v <- 1 - runif(1) # as exceedance, in order to reinforce the
# change to exceedance instead of nonexceedance that otherwise dominates this package
message("Exceedance probabilities u' and v' are ", u, " and ", v)
coCOP(u,v,cop=PLACKETTcop, para=10) # Positive association Plackett

# computation using manual manipulation to nonexceedance probability
1 - COP(cop=PSP,(1-u),(1-v))
# computation using internal manipulation to nonexceedance probability
coCOP(cop=PSP, u, v)

# Next demonstrate COP + coCOP = unity.
"MOcop.formula" <- function(u,v, para=para, ...) { # Marshall-Olkin copula
  alpha <- para[1]; beta <- para[2]; return(min(v*u^(1-alpha), u*v^(1-beta)))}
```

```

}
"M0cop" <- function(u,v, ...) { asCOP( u, v, f=M0cop.formula, ...) }
u <- 0.2; v <- 0.75; ab <- c(1.5, 0.3)
COP(u,v, cop=M0cop, para=ab) + coCOP(1-u,1-v, cop=M0cop, para=ab) # UNITY

```

composite1COP

Composition of a Single Symmetric Copula with Two Compositing Parameters (Khoudraji Device with Pi Independence)

Description

The *composition of a single copula* (Salvadori *et al.*, 2006, p. 266, prop. C.3) is created by the following result related to “composition of copulas” in that reference. This construction technique is named the *Khoudraji device* within the **copula** package (see `khoudrajiCopula` therein) (Hofert *et al.*, 2018, pp. 120–121). Suppose $\mathbf{C}(u, v)$ is a *symmetric copula* (see [COP](#)) with parameters Θ and $\mathbf{C} \neq \mathbf{\Pi}$ (for $\mathbf{\Pi}$ see [P](#)), then a family of generally *asymmetric copulas* $\mathbf{C}_{\alpha, \beta; \Theta}$ with **two compositing parameters** $0 < \alpha, \beta < 1$, and $\alpha \neq \beta$, which also includes just the copula $\mathbf{C}(u, v)$ as a limiting case for $\alpha = \beta = 0$ and is given by

$$\mathbf{C}_{\alpha, \beta}(u, v) = u^{\alpha} v^{\beta} \cdot \mathbf{C}(u^{1-\alpha}, v^{1-\beta}).$$

The `composite1COP` function provides the means for inserting *permutation asymmetry* from a *permutation symmetric* copula as described by Joe (2017, p. 124), but do so in a more general way through the provision of two and not just one parameter. Joe’s description is supported herein if one of the α or β is held at zero. Very loosely, the $\alpha > 0$ kicks probability density down towards the lower right corner, whereas $\beta > 0$ kicks density up towards the upper left corner. Finally, the `composite2COP` function is based on a slightly more general result that is shown with `composite2COP`, which provides further details of copula composition and more contextualization of Hofert *et al.* (2018) remarks on the *Khoudraji device*.

Usage

```

composite1COP(u, v, para, ...)
khoudraji1COP(u, v, para, ...)
khoudrajiPCOP(u, v, para, ...)

```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A special parameter list (see Note); and
<code>...</code>	Additional arguments to pass to the copula.

Value

Value(s) for the composited copula are returned.

Note

The following descriptions list in detail the structure and content of the para argument:

alpha — The α compositing parameter;
 beta — The β compositing parameter;
 cop1 — Function of the copula $C(u, v)$; and
 para1 — Vector of parameters Θ_C for $C(u, v)$.

For the para argument, the same nomenclature as used for [composite2COP](#) is used with obviously cop2 and para2 dropped for composite1COP. The cop1 and para1 names remain enumerated for composite1COP so that the para argument of the more general [composite2COP](#) function could be used directly in [composite1COP](#). Albeit, the second copula and its parameters would not be used. A more complex (extended) composition in [composite3COP](#) extends this basic parameter structure.

Author(s)

W.H. Asquith

References

Hofert, M., Kojadinovic, I., Mächler, M., and Yan, J., 2018, Elements of copula modeling with R: Dordrecht, Netherlands, Springer.

Joe, H., 2017, Parametric copula families for statistical models (chap. 8) in Copulas and dependence models with applications—Contributions in honor of Roger B. Nelsen, *eds.* Flores, U.M., Amo Artero, E., Durante, F., Sánchez, J.F.: Springer, Cham, Switzerland, ISBN 978–3–319–64220–9, [doi:10.1007/9783319642215](#).

Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, Extremes in Nature—An approach using copulas: Springer, 289 p.

See Also

[COP](#), [breveCOP](#), [composite2COP](#), [composite3COP](#), [convexCOP](#), [glueCOP](#)

Examples

```
## Not run:
alpha <- 0.24; beta <- 0.23; Theta1 <- NA;
# W() does not use parameters, but show how parameters would be set if needed.
para <- list(alpha=alpha, beta=beta, cop1=W, para1=Theta1)
t <- composite1COP(0.4, 0.6, para)
if( t != W(0.4, 0.6)) message("Not equal as expected") #
## End(Not run)

## Not run:
# Hofert et al. (2018, p. 124, eq. 3.15)
# No matter what copula is chosen, Kendall tau must be
# Tau <= (alpha * beta) / (alpha + beta - alpha * beta)
# and those authors report Tau <= 0.5816. We can test this computation by
para <- list(cop=M, para=NULL, alpha=1-0.6, beta=1-0.95)
```

```

    tauCOP(khoudrajiPCOP, para=para) # 0.5816283
## End(Not run)

## Not run:
# Next use this as a chance to check logic flow through the various
# "compositing" operators and their use as needed dispatch to COP().
my.para <- list(cop1=GHcop, para1=exp(+1.098612) + 1,
               cop2=PLcop, para2=exp(-1.203973),
               alpha=0.5, beta=0.25, kappa=0.1, gamma=0.1,
               weights=c(0.95, 0.05))
# uses cop1/2, para1/2, only weights
nustarCOP(cop=convexCOP, para=my.para) # 0.8570434

# uses cop1/2, para1/2, only alpha
nustarCOP(cop=convex2COP, para=my.para) # 0.2697063

# uses cop1, para1, only alpha / beta
nustarCOP(cop=composite1COP, para=my.para) # 0.5103119

# uses cop1/2, para1/2, only alpha / beta
nustarCOP(cop=composite2COP, para=my.para) # 0.0714571

# uses cop1/2, para1/2, only alpha, beta, kappa, gamma
nustarCOP(cop=composite3COP, para=my.para) # 0.0792634
## End(Not run)

## Not run:
# Hofert et al. (2018, p. 121, fig. 3.20, left panel)
# The ordering of copula and the "1-" operations on alpha and beta in copBasic
# differ from that shown in Hofert et al. (2018), but instead of their
# "kho(0.6, 0.95)(CLcop(6), P)" notation for the their left panel, in copBasic
# we can reproduce their simulation by the following. So, swapping notation
# between the copula package (khoudrajiCopula) and copBasic would be required.
para <- list(cop=CLcop, para=6, alpha=1-0.95, beta=1-0.6, reflect=1)
UV <- simCOP(n=5000, cop=khoudrajiPCOP, para=para) # then do it again with
# reflect = 2, then 3, then 4 to show reflection/rotation dispatch to COP().
## End(Not run)

```

composite2COP

*Composition of Two Copulas with Two Compositing Parameters
(Khoudraji Device)*

Description

The *composition of two copulas* (Salvadori *et al.*, 2007, p. 266, prop. C.3) provides for more sophisticated structures of dependence between variables than many single parameter copula can provide. Further, *asymmetrical copulas* are readily obtained from *symmetrical copulas*. Let **A** and **B** be copulas with respective parameters Θ_A and Θ_B , then

$$\mathbf{C}_{\alpha,\beta}(u, v) = \mathbf{A}(u^\alpha, v^\beta) \cdot \mathbf{B}(u^{1-\alpha}, v^{1-\beta}),$$

defines a family of copulas $\mathbf{C}_{\alpha,\beta;\Theta_A,\Theta_B}$ with **two** *compositing parameters* $\alpha, \beta \in \mathcal{I} : [0, 1]$. In particular if $\alpha = \beta = 1$, then $\mathbf{C}_{1,1} = \mathbf{A}$, and if $\alpha = \beta = 0$, then $\mathbf{C}_{0,0} = \mathbf{B}$. For $\alpha \neq \beta$, the $\mathbf{C}_{\alpha,\beta}$ is in general asymmetric that is $\mathbf{C}(u, v) \neq \mathbf{C}(v, u)$ for some $(u, v) \in \mathcal{I}^2$. This construction technique is named the *Khoudraji device* within the **copula** package (see `khoudrajiCopula` therein) (Hofert *et al.*, 2018, p. 120).

It is important to stress that copulas $\mathbf{A}_{\Theta_A}$ and $\mathbf{B}_{\Theta_B}$ can be of different families and each copula parameterized accordingly by the vector of parameters Θ_A and Θ_B . This is an interesting feature in the context of building complex structures when pursuing asymmetric measures of dependency such as the *L-comoments*. Symmetry of the copula $\mathbf{C}$ is required for the situation that follows, however.

It is possible to simplify the construction of an asymmetric copula from a symmetric copula by the following. Let $\mathbf{C}(u, v)$ be a symmetric copula, $\mathbf{C} \neq \mathbf{\Pi}$ (for $\mathbf{\Pi}$ see [P](#)). A family of asymmetric copulas $\mathbf{C}_{\alpha,\beta}$ with **two** *composition parameters* $0 < \alpha, \beta < 1$, and $\alpha \neq \beta$ that also includes $\mathbf{C}(u, v)$ as a limiting case and is given by

$$\mathbf{C}_{\alpha,\beta}(u, v) = u^\alpha v^\beta \cdot \mathbf{C}(u^{1-\alpha}, v^{1-\beta}).$$

Hofert *et al.* (2018, p. 121) comment that “from a practical perspective, a useful subset of families constructed from [the] Khoudraji device is obtained” by choosing *independence copula* ($\mathbf{\Pi}$, [P](#)) for one of the copula and that choosing $[\alpha$ or $\beta]$ relative close to 1 produces nonexchangeable [see `isCOP.permsym`] versions of $\mathbf{C}(u, v)$ (meaning $\mathbf{C}(u, v) \neq \mathbf{C}(v, u)$). For the **copBasic** package, the `khoudraji1COP()` and `khoudrajiPCOP()` (the P meaning [P](#)) aliases are added for programming clarity for developers desiring to have contrasting copula calls to the three *compositing* copula: `composite1COP`, `composite2COP()`, and `composite3COP`.

Usage

```
composite2COP(u, v, para, ...)
khoudraji2COP(u, v, para, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A special parameter list (see Note); and
<code>...</code>	Additional arguments to pass to the copulas.

Value

Value(s) for the composited copula is returned.

Note

The following descriptions list in detail the structure and content of the `para` argument:

`alpha` — The α compositing parameter;
`beta` — The β compositing parameter;
`cop1` — Function of the first copula **A**;
`cop2` — Function of the second copula **B**;
`para1` — Vector of parameters Θ_A for **A**; and
`para2` — Vector of parameters Θ_B for **B**.

The `para` argument of this function also can be passed to [composite1COP](#); albeit, the second copula and its parameters would not be used. A more complex (extended) composition in [composite3COP](#) extends this basic parameter structure.

Author(s)

W.H. Asquith

References

Hofert, M., Kojadinovic, I., Mächler, M., and Yan, J., 2018, Elements of copula modeling with R: Dordrecht, Netherlands, Springer.

Salvadori, G., De Michele, C., Kotegoda, N.T., and Rosso, R., 2007, Extremes in Nature—An approach using copulas: Springer, 289 p.

See Also

[COP](#), [breveCOP](#), [composite1COP](#), [composite3COP](#), [convexCOP](#), [glueCOP](#)

Examples

```
alpha <- 0.24; beta <- 0.23; Theta1 <- NA; Theta2 <- NA
# The W() and PSP() copulas do not take parameters, but example shows how the
# parameters would be set should either or both of the copulas require parameters.
para <- list(alpha=alpha, beta=beta, cop1=W, cop2=PSP, para1=Theta1, para2=Theta2)
print(composite2COP(0.4, 0.6, para)) # 0.2779868

# In this example, the N4212cop uses "3" as its parameter value.
para <- list(alpha=alpha, beta=beta, cop1=W, cop2=N4212cop, para1=Theta1, para2=3)
print(composite2COP(0.4, 0.6, para)) # 0.3387506

## Not run:
# This example does a great job of showing a composited copula with a near singularity,
# but with leakage of chance to the upper left. The example is also critical because
# it shows that gridCOP is returning a matrix in the proper orientation relative to
# the level.curvesCOP and simCOP functions. Example is cross-ref'ed from gridCOP() docs.
layout(matrix(1:2,byrow=TRUE))
para <- list(alpha=0.5, beta=0.90, cop1=M, cop2=N4212cop, para1=NA, para2=1.4)
image(gridCOP(cop=composite2COP, para=para, delta=0.01), col=terrain.colors(30),
```

```

      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
D <- simCOP(n=2000, cop=composite2COP, para=para, ploton=FALSE, pch=4, col=4, cex=0.75)
level.curvesCOP(cop=composite2COP, para=para, ploton=FALSE, delt=0.05)
mtext("Theoretical composited copula, level curves, and simulation")

emp <- EMPIRgrid(para=D, deluv=0.05)      # CPU heavy
image(emp$empcop, col=terrain.colors(30) ) # image orientation is correct!
# Depending on balance between sample size, deluv, delu, and delt, one or more:
# Error in uniroot(func, interval = c(0, 1), u = u, LHS = t, cop = cop, :
#   f() values at end points not of opposite sign
# warnings might be triggered. This is particularly true because of the flat derivative
# above the near singularity in this example composited copula.
points(D$U, D$V, pch=4, col="blue", cex=0.75)
level.curvesCOP(cop=EMPIRcop, para=D, ploton=FALSE, delu=0.02, delt=0.05)
mtext("Empirical copula from n=2000 simulation") #
## End(Not run)

## Not run:
# Demonstration in how to add reflection (rotation by 180) degrees of the GHcop
# and use the two-parameter version of that coupla to accent upper tail coupling
# with a much weaker lower tail, but we want open up or spread the middle part of
# the distribution. Lastly, because alpha != beta, permutation asymmetry is added.
para <- list(cop1=W, para1=NA, cop2=GHcop, para2=c(1,10), alpha=0.01, beta=0.06)
UV <- simCOP(1000, cop=COP, para=list(cop=composite2COP, reflect=2, para=para)) #
## End(Not run)

```

composite3COP

(Extended) Composition of Two Copulas with Four Compositing Parameters

Description

The *(extended) composition of two copulas* (Salvadori *et al.*, 2006, p. 266, prop. C.4) provides for even more sophisticated structures of dependence between variables than two-copula composition in [composite2COP](#). Let **A** and **B** be copulas with respective parameters Θ_A and Θ_B , then

$$\mathbf{C}_{\alpha, \beta, \kappa, \gamma}(u, v) = u^{\kappa} v^{\gamma} \cdot \mathbf{A}([u^{1-\kappa}]^{\alpha}, [v^{1-\gamma}]^{\beta}) \cdot \mathbf{B}([u^{1-\kappa}]^{1-\alpha}, [v^{1-\gamma}]^{1-\beta}),$$

defines a family of copulas $\mathbf{C}_{\alpha, \beta, \kappa, \gamma}$ with **four** *compositing parameters* $\alpha, \beta, \kappa, \gamma \in (0, 1)$.

It is important to stress that copulas $\mathbf{A}_{\Theta_A}$ and $\mathbf{B}_{\Theta_B}$ can be of different families and each parameterized accordingly by the vectors of parameters Θ_A and Θ_B .

Usage

```
composite3COP(u, v, para, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in X direction;
<code>v</code>	Nonexceedance probability v in Y direction;
<code>para</code>	A special parameter list (see Note); and
<code>...</code>	Additional arguments to pass to composite2COP .

Value

A value for the composited copula is returned.

Note

The following descriptions list in detail the structure and content of the `para` argument:

- `alpha` — The α compositing parameter;
- `beta` — The β compositing parameter;
- `kappa` — The κ compositing parameter;
- `gamma` — The γ compositing parameter;
- `cop1` — Function of the first copula **A**;
- `cop2` — Function of the second copula **B**;
- `para1` — Vector of parameters $\Theta_{\mathbf{A}}$ for **A**; and
- `para2` — Vector of parameters $\Theta_{\mathbf{B}}$ for **B**.

The first example produces two plots. These are extremely informative for many nuances of copula theory. Whereas it is difficult in prose to describe, users are strongly encouraged that once full understanding of connection of red and green between the easier to understand bivariate plot and the plot showing the sections and derivatives of the sections is achieved that much of copula theory will be mastered—get a copy of Nelsen (2006) and (or) Salvadori *et al.* (2007).

Author(s)

W.H. Asquith

References

- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, Extremes in Nature—An approach using copulas: Springer, 289 p.

See Also

[COP](#), [breveCOP](#), [simCOP](#), [composite1COP](#), [composite2COP](#), [convexCOP](#), [glueCOP](#), [simcomposite3COP](#)

Examples

```
## Not run:
para <- list(cop1=PLACKETTcop, cop2=N4212cop,
             para1=10^(runif(1,min=-5,max=5)), para2=runif(1,min=1,max=100),
             alpha=runif(1), beta=runif(1), kappa=runif(1), gamma=runif(1))
txts <- c("Alpha=", round(para$alpha, digits=4),
          "; Beta=", round(para$beta, digits=4),
          "; Kappa=", round(para$kappa, digits=4),
          "; Gamma=", round(para$gamma, digits=4),
          "; Theta1=", round(para$para1[1], digits=5),
          "; Theta2=", round(para$para2[1], digits=2))
layout(matrix(1:2, byrow=TRUE))
D <- simCOP(n=300, cop=composite3COP, para=para, cex=0.5, col=rgb(0,0,0.2), pch=16)
mtext(paste(txts,collapse=""))

f <- round(runif(1),digits=2)
ftxt <- c("Sectionals (thick) and derivatives (thin) at f=",f," nonexceedance prob.")
segments(f,0,f,1, col=3, lwd=2); segments(0,f,1,f, col=2, lwd=2)
t <- sectionCOP(f,cop=composite3COP,para=para, col=3, lwd=4)
t <- sectionCOP(f,cop=composite3COP,para=para, dercop=TRUE, ploton=FALSE,col=3)
t <- sectionCOP(f,cop=composite3COP,para=para, wrtV=TRUE, ploton=FALSE,col=2,lwd=4)
t <- sectionCOP(f,cop=composite3COP,para=para, wrtV=TRUE, ploton=FALSE,col=2,
               dercop=TRUE)
mtext(paste(ftxt, collapse=""))#
## End(Not run)
```

convex2COP

Convex Combination of Two Copulas

Description

The *convex composition of two copulas* (Joe, 2014, p. 155) provides for some simple complexity extension between copula families. Let **A** and **B** be copulas with respective vectors of parameters Θ_A and Θ_B , then the convex combination of these copulas is

$$C_{\alpha}^{\times}(u, v) = \alpha \cdot \mathbf{A}(u, v; \Theta_A) - (1 - \alpha) \cdot \mathbf{B}(u, v; \Theta_B),$$

where $0 \leq \alpha \leq 1$. The generalization of this function for N number of copulas is provided by [convexCOP](#).

Usage

```
convex2COP(u,v, para, ...)
```

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction;
para	A special parameter list (see Note); and
...	Additional arguments to pass to the copula.

Value

Value(s) for the convex combination copula is returned.

Note

The following descriptions list in detail the structure and content of the para argument:

alpha — The α compositing parameter;
 cop1 — Function of the first copula **A**;
 cop2 — Function of the second copula **B**;
 para1 — Vector of parameters $\Theta_{\mathbf{A}}$ for **A**; and
 para2 — Vector of parameters $\Theta_{\mathbf{B}}$ for **B**.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[COP](#), [breveCOP](#), [convexCOP](#), [composite1COP](#), [composite2COP](#), [composite3COP](#), [FRECHETcop](#), [glueCOP](#)

Examples

```
para <- list(alpha=0.24, cop1=FRECHETcop, para1=c(0.4, 0.56),
             cop2=PSP, para2=NA)
convex2COP(0.87,0.35, para=para) # 0.3188711
## Not run:
# Suppose we have a target Kendall Tau of 1/3 and a Gumbel-Hougaard copula seems
# attractive but the GH has just too much upper tail dependency for comfort. We
# think from data analysis that an upper tail dependency that is weaker and near
# 2/10 is the better. Let us convex mix in a Plackett copula and optimize.
TargetTau <- tauCOP(cop=GHCop, para=1.5) # 1/3 (Kendall Tau)
taildepCOP( cop=GHCop, para=1.5, plot = TRUE)$lambdaU # 0.4126
TargetUpperTailDep <- 2/10

# **Serious CPU time pending for this example**
par <- c(-.10, 4.65) # Initial guess but the first parameter is in standard
# normal for optim() to keep us in the [0,1] domain when converted to probability.
# The guesses of -.10 (standard deviation) for the convex parameter and 4.65 for
# the Plackett are based on a much longer search times as setup for this problem.
# The simplex for optim() is going to be close to the solution on startup.
"afunc" <- function(par) {
  para <- list(alpha=pnorm(par[1]), cop1=GHCop, para1=1.5,
               cop2=PLACKETTcop, para2=par[2])
  tau <- tauCOP(cop=convex2COP, para=para)
```

```

    taildep <- taildepCOP(cop=convex2COP, para=para, plot = FALSE)$lambdaU
    err <- sqrt((TargetTau - tau)^2 + (TargetUpperTailDep - taildep)^2)
    print(c(pnorm(par[1]), par[2], tau, taildep, err))
    return(err)
}
mysolution <- optim(par, afunc, control=list(abstol=1E-4))

para <- list(alpha=.4846902, cop1=GHcop,      para1=1.5,
             cop2=PLACKETTcop, para2=4.711464)
UV <- simCOP(n=2500, cop=convex2COP, para=para, snv=TRUE) #
## End(Not run)

```

convexCOP

*Convex Combination of an Arbitrary Number of Copulas***Description**

The *convex composition of N number of copulas* (Salvadori *et al.*, p. 132, 2007) provides for complexity extension between coupla families. Let $\mathbf{C}_i$ be a copula with respective vector of parameters Θ_i , then the convex combination of these copulas is

$$\mathbf{C}_{\omega}^{\times}(u, v) = \sum_{i=1}^N \omega_i \mathbf{C}_i(u, v; \Theta_i),$$

where $\sum_{i=1}^N \omega_i = 1$ for N number of copulas. The weights ω are silently treated as $1/N$ if the weights element is absent in the R list argument para.

Usage

```
convexCOP(u,v, para, ...)
```

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction;
para	A special parameter list (see Note); and
...	Additional arguments to pass to the copula.

Value

Value(s) for the convex combination copula is returned.

Note

The following descriptions list in detail the structure and content of the `para` argument but please reference the **Examples** to see the `i` notation:

`copi` — The i th copula;

`parai` — Vector of parameters Θ_i ; and

`weights` — Optional vector of weights whose sum will be rescaled to unity; default is $1/N$ for each weight.

Author(s)

W.H. Asquith

References

Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, *Extremes in Nature—An approach using copulas*: Springer, 289 p.

See Also

[COP](#), [breveCOP](#), [convex2COP](#), [composite1COP](#), [composite2COP](#), [composite3COP](#), [glueCOP](#)

Examples

```
# The copulas and parameters are named by sequence number appended to cop and para.
para1 <- list(cop1=GHcop, cop2=PLcop, para1=8, para2=.03, weights=c(.8,.2))
para2 <- list(cop1=GHcop, cop2=PLcop, para1=8, para2=.03, alpha=0.8)
H <- convexCOP( 0.6,0.4, para=para1)
G <- convex2COP(0.6,0.4, para=para2)
if( abs(H-G) <= 1e-6 ) message("They are equal.")

## Not run:
# A convex combination of three copulas. A GHcop with strong positive association and
# a Plackett with strong negative association, and independence. The weights favor the
# GHcop but a little outlier and expansive spread is superimposed on the core trend.
para <- list(cop1=GHcop, cop2=PLcop, cop3=P,
             para1=8, para2=.03, para3=NA, weights=c(40,7,10))
UV <- simCOP(1000, cop=convexCOP, para=para, lwd=0.8) #
## End(Not run)
```

Description

Compute the *copula* or *joint distribution function* through a copula as shown by Nelsen (2006, p. 18) is the joint probability

$$\Pr[U \leq u, V \leq v] = \mathbf{C}(u, v).$$

The copula is an expression of the joint probability that both $U \leq u$ and $V \leq v$.

A copula is a type of *dependence function* that permits straightforward characterization of dependence from independence. Joe (2014, p. 8) comments that “copula families are usually given as cdfs [cumulative distribution functions.]” A *radially symmetric* or *permutation symmetric copula* is one such that $\mathbf{C}(u, v) = \mathbf{C}(v, u)$ otherwise the copula is *asymmetric*.

The copula *inversions* $t = \mathbf{C}(u=U, v)$ or $t = \mathbf{C}(u, v=V)$ for a given t and U or V are provided by [COPinv](#) and [COPinv2](#), respectively. A copula exists in the domain of the unit square ($\mathcal{I}^2 = [0, 1] \times [0, 1]$) and is a *grounded* function meaning that

$$\mathbf{C}(u, 0) = 0 = \mathbf{C}(0, v) \text{ and thus } \mathbf{C}(0, 0) = 0,$$

and other properties of a copula are that

$$\mathbf{C}(u, 1) = u \text{ and } \mathbf{C}(1, v) = v \text{ and}$$

$$\mathbf{C}(1, 1) = 1.$$

Copulas can be combined with each other ([convexCOP](#), [convex2COP](#), [composite1COP](#), [composite2COP](#), [composite3COP](#), and [glueCOP](#)) to form more complex and sophisticated dependence structures. Also copula multiplication—a special product of two copulas—yields another copula (see [prod2COP](#)).

Perhaps the one of the more useful features of this function is that in practical applications it can be used to take a copula formula and reflect or rotated it in fashions to attain association structures that the native definition of the copula can not acquire. The terminal demonstration in the **Examples** demonstrates this for the *Raftery copula* ([RFCop](#)).

Usage

```
COP(u, v, cop=NULL, para=NULL,
    reflect=c("cop", "surv", "acute", "grave",
              "1", "2", "3", "4"), ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>cop</code>	A copula function with vectorization as in <code>asCOP</code> ;
<code>para</code>	Vector of parameters or other data structures, if needed, to pass to the copula;
<code>reflect</code>	The reflection of the copula form (see Note) and the default “cop” or “1” is the usual copula definition (also see simCOPmicro). The numbered values correspond, respectively, to the named values; and
<code>...</code>	Additional arguments to pass to the copula.

Value

Value(s) for the copula are returned.

Note

REFLECTIONS OF VARIABLES (ROTATIONS OF THE COPULA)—The copula of $(1 - U, 1 - V)$ is the survival copula $\hat{\mathbf{C}}(u, v)$; [surCOP](#) and is defined as

$$\Pr[U > u, V > v] = \hat{\mathbf{C}}(u, v) = u + v - 1 + \mathbf{C}(1 - u, 1 - v) \rightarrow \text{"surv"},$$

whereas, following the notation of Joe (2014, p. 271), the copula of $(1 - U, V)$ is defined as

$$\Pr[U > u, V \leq v] = \acute{\mathbf{C}}(u, v) = v - \mathbf{C}(1 - u, v) \rightarrow \text{"acute"}, \text{ and}$$

the copula of $(U, 1 - V)$ is defined as

$$\Pr[U \leq u, V > v] = \grave{\mathbf{C}}(u, v) = u - \mathbf{C}(u, 1 - v) \rightarrow \text{"grave"}.$$

Here it is useful to stress the probability aspects that change with the reflections, but this section ends with the reflections themselves being graphically highlighted. The **Examples** stress simple variations on the probability aspects.

To clarify the seemingly clunky nomenclature—Joe (2014) does not provide “names” for $\acute{\mathbf{C}}(u, v)$ or $\grave{\mathbf{C}}(u, v)$ —the following guidance is informative where the numbers in the list align to those for the reflect argument:

- (2) "surv" or $\hat{\mathbf{C}}(u, v)$ is a reflection of U and V on the horizontal *and* vertical axes, respectively,
- (3) "acute" or $\acute{\mathbf{C}}(u, v)$ is a reflection of U on the horizontal axis, and
- (4) "grave" or $\grave{\mathbf{C}}(u, v)$ is a reflection of V on the vertical axis.

The names "acute" and "grave" match those used in the **Rd**-format math typesetting instructions. Users are directed to the documentation of [simCOPmicro](#) for further discussion because the COP function is expected to be an early entry point for new users interested in the **copBasic** API.

For the **copBasic** package and in order to keep some logic brief and code accessible for teaching and applied circumstances, reflections of copulas using analogs to the reflect argument are only natively supported in the COP and [simCOPmicro](#) functions. The interfaces of **copBasic** should already be flexible enough for users to adapt and (or) specially name reflections of copulas for deployment. A caveat is that some individual copula implementations might have some self-supporting infrastructure. The reflection can also be set within the para argument when it is a list (see **Examples**).

An example is warranted. Although the Gumbel–Hougaard copula ([GHcop](#)) can be reflected by COP and [simCOPmicro](#) and testing is made in the **Note** section of [simCOPmicro](#), it is suggested that a user generally requiring say a horizontal reflection ru (or vertical reflection rv) of the Gumbel–Hougaard copula write a function named perhaps ruGHcop (or rvGHcop).

Such functions, consistent with the mathematics at the beginning of this **Note**, can be used throughout functions of **copBasic** using the cop arguments. The author (Asquith) eschews implementing what is perceived as too much flexibility and overhead for the package to support the three reflection permutations universally across all copula functions of the package. This being said, COP can take an R list for the para argument for rotation/reflection:

```
set.seed(14)
UV3 <- simCOP(20, cop=COP, pch=16, col=3,
```

```

      para=list(cop=GLcop, para=pi+1, reflect="3"))
set.seed(14)
UV2 <- simCOP(20, cop=COP, pch=16, col=4, ploton=FALSE,
      para=list(cop=GLcop, para=pi+1, reflect="2"))
arrows(x0=UV3[,1], y0=UV3[,2], x=UV2[,1], y=UV2[,2])

```

and this type of interface is similar to [composite1COP](#) as the following rotation and then asymmetric construction shows:

```

UV <- simCOP(1000, cop=composite1COP,
      para=list(cop1=COP,
      para1=c(cop=GHCop, para=pi+1, reflect="4"),
      alpha=0.1, beta=0.3))

```

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
 Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[coCOP](#), [duCOP](#), [surCOP](#), [surfuncCOP](#)

Examples

```

u <- runif(1); v <- runif(1)
COP(cop=W,u,v); COP(cop=P,u,v); COP(cop=M,u,v); COP(cop=PSP,u,v)

FF <- 0.75 # 75th percentile, nonexceedance
GG <- 0.20 # 25th percentile, nonexceedance
bF <- 1 - FF; bG <- 1 - GG # exceedance
# What is the probability that both X and Y are less than
# 75th and 20th percentiles, respectively?
COP(cop=P, FF, GG) # 0.15
# What is the probability that both X and Y are greater than
# 75th and 20th percentiles, respectively?
surCOP(cop=P, bF, bG) # 0.20
# What is the probability that either X or Y are less than
# the 75th and 20th percentiles, respectively?
duCOP(cop=P, FF, GG) # 0.8
# What is the probability that either X or Y are greater than
# the 75th and 20th percentiles, respectively?
coCOP(cop=P, bF, bG) # 0.85

# Repeat for the PSP copula:
# What is the probability that both X and Y are less than
# 75th and 20th percentiles, respectively?

```

```

COP(cop=PSP, FF, GG) # 0.1875
# What is the probability that both X and Y are greater than
# 75th and 20th percentiles, respectively?
surCOP(cop=PSP, bF, bG) # 0.2375
# What is the probability that either X or Y are less than
# the 75th and 20th percentiles, respectively?
duCOP(cop=PSP, FF, GG) # 0.7625
# What is the probability that either X or Y are greater than
# the 75th and 20th percentiles, respectively?
coCOP(cop=PSP, bF, bG) # 0.8125
# Both of these summations equal unity
  COP(cop=PSP, FF, GG) + coCOP(cop=PSP, bF, bG) # 1
surCOP(cop=PSP, bF, bG) + duCOP(cop=PSP, FF, GG) # 1

FF <- 0.99 # 99th percentile, nonexceedance
GG <- 0.50 # 50th percentile, nonexceedance
bF <- 1 - FF # nonexceedance
bG <- 1 - GG # nonexceedance
# What is the probability that both X and Y are less than
# 99th and 50th percentiles, respectively?
COP(cop=P, FF, GG) # 0.495
# What is the probability that both X and Y are greater than
# 99th and 50th percentiles, respectively?
surCOP(cop=P, bF, bG) # 0.005
# What is the probability that either X or Y are less than
# the 99th and 50th percentiles, respectively?
duCOP(cop=P, FF, GG) # 0.995
# What is the probability that either X or Y are greater than
# the 99th and 50th percentiles, respectively?
coCOP(cop=P, bF, bG) # 0.505

## Not run:
# MAJOR EXAMPLE FOR QUICKLY MODIFYING INHERENT ASSOCIATION STRUCTURES
p <- 0.5 # Reasonable strong positive association for the Raftery copula
"RFcop1" <- function(u,v, para) COP(u,v, cop=RFcop, para=para, reflect="1")
"RFcop2" <- function(u,v, para) COP(u,v, cop=RFcop, para=para, reflect="2")
"RFcop3" <- function(u,v, para) COP(u,v, cop=RFcop, para=para, reflect="3")
"RFcop4" <- function(u,v, para) COP(u,v, cop=RFcop, para=para, reflect="4")

d <- 0.01 # Just to speed up the density plots a bit
densityCOPplot(RFcop1, para=p, contour.col=1, deluv=d) # Raftery in the literature
densityCOPplot(RFcop2, para=p, contour.col=1, deluv=d, ploton=FALSE)
densityCOPplot(RFcop3, para=p, contour.col=1, deluv=d, ploton=FALSE)
densityCOPplot(RFcop4, para=p, contour.col=1, deluv=d, ploton=FALSE)
# Now some text into the converging tail to show the reflection used.
text(-2,-2, "reflect=1", col=2); text(+2,+2, "reflect=2", col=2)
text(+2,-2, "reflect=3", col=2); text(-2,+2, "reflect=4", col=2) #
## End(Not run)

## Not run:
# ALTERNATIVE EXAMPLE FOR QUICKLY MODIFYING INHERENT ASSOCIATION STRUCTURES
# To show how the reflection can be alternatively specified and avoid in this case
# making four Raftery functions, pass by a list para argument. Also, demonstrate

```

```

# that cop1 --> cop and para1 --> para are the same in use of the function. This
# provides some nomenclature parallel to the other compositing functions.
densityCOPplot(COP, para=list(reflect=1, cop1=RFcop, para=p ), deluv=d,
               contour.col=1, drawlabels=FALSE)
densityCOPplot(COP, para=list(reflect=2, cop= RFcop, para1=p), deluv=d,
               contour.col=2, drawlabels=FALSE, ploton=FALSE)
densityCOPplot(COP, para=list(reflect=3, cop1=RFcop, para1=p), deluv=d,
               contour.col=3, drawlabels=FALSE, ploton=FALSE)
densityCOPplot(COP, para=list(reflect=4, cop= RFcop, para=p ), deluv=d,
               contour.col=4, drawlabels=FALSE, ploton=FALSE)
# Now some text into the converging tail to show the reflection used.
text(-2,-2, "reflect=1", col=2); text(+2,+2, "reflect=2", col=2)
text(+2,-2, "reflect=3", col=2); text(-2,+2, "reflect=4", col=2) #
## End(Not run)

## Not run:
# Similar example to previous, but COP() can handle the reflection within a
# parameter list ,and the reflect, being numeric here, is converted to
# character internally.
T12 <- CLcop(tau=0.67)$para # Kendall Tau of 0.67
T12 <- list(cop=CLcop, para=T12, reflect=2) # reflected to upper tail dependency
UV  <- simCOP(n=1000, cop=COP, para=T12) #
## End(Not run)

```

copBasic.fitpara

A Single or Multi-Parameter Optimization Engine (Beta Version)

Description

An example of a general implementation of a parameter optimization scheme using core features of the **copBasic** package. Because of the general complexity of the objectives for this function, the interface shown here is considered an “beta version” and nomenclature is subject to possibly sweeping changes in the future.

Usage

```
copBasic.fitpara.beta(uv=NULL, popstat=NULL, statf=NULL, cop=NULL,
                     paradim=1, interval=NULL, par.init=NULL, ...)
```

Arguments

uv	An R two column matrix or data.frame of a sample of nonexceedance probabilities u and v ;
popstat	The population statistic(s) that will be used if uv is NULL;
statf	A function responsible at the minimum for computation of the theoretical values of the population statistic(s) that the optimization will revolve around; This function, if supporting an as.sample interface (e.g. hoefCOP) and if uv has been provided, will be dispatched to compute the population statistic(s);

cop	A copula function that is passed along to statf though of course the statf function can decide whether to use this argument or not;
paradim	The dimension of the parameters. In reality, the default triggers uni-dimensional root solving using the uniroot() function in R or otherwise the optim() function in R is used for multi-dimensional minimization with subtle changes in setup (see source code). Alternative logic could be have been used but it is felt that this is the most logical for future adaptations;
interval	The interval argument by the same name for the uniroot() function;
par.init	The initial parameter vector for the optim() function; and
...	Additional arguments to pass.

Value

A vector of the values for the parameter variable is returned

Note

One-Parameter Optimization—A demonstration of one-dimensional parameter optimization using the *Gini Gamma* ([giniCOP](#)), which is a measure of bivariate association. There is no native support for Gini Gamma (and most of the other measures of association) in regards to being a parameter estimator at the copula function interface level in **copBasic**. (A converse example is one provided internally by the [GHcop](#) copula.)

```
set.seed(24); n <- 230 # sample size to draw from Galambos copula but a
# different copula was chosen for the fitting.
sampleUV <- simCOP(n=n, cop=GLcop, para=1.5) # a random sample
para <- copBasic.fitpara.beta(uv=sampleUV, statf=giniCOP,
                             interval=c(.1,200), cop=HRcop) # 1.959521
```

Three-Parameter Optimization—A demonstration of multi-dimensional parameter optimization using the *Gini Gamma* ([giniCOP](#)), *Nu-Skew* ([nuskewCOP](#)), and *Nu-Star* ([nustarCOP](#)). This is substantially more complicated to implement. The *Hüsler-Reiss copula* ([HRcop](#)) is chosen both as part of the sample simulation for the study as well as the copula as part of the modeling. Using composition by the [composite1COP](#), first establish a parent three parameter copula and simulate from it to create a bivariate sample in sampleUV that will be used for demonstration. A standard normal variate graphic of the simulation is generated by [simCOP](#) as well—later, additional results will be superimposed.

```
n <- 610; set.seed(1999) # Seed value will be used again (see below)
pop.para <- list(cop1=HRcop, para1=4, alpha=0.14, beta=0.35)
sampleUV <- simCOP(n=n, cop=composite1COP, para=pop.para, col=3, snv=TRUE)
```

A custom objective function objfunc to serve as the statf for the copBasic.fitpara.beta call. The objective function has the as.sample interface (e.g. [giniCOP](#)) implemented for sample estimation. The most subtle feature of function presumably is the use of the pnorm() function in R for the α and β parameters to keep each parameter in the range $\alpha, \beta \in (0, 1)$. Another subtly, which affects the choice of other copulas from [HRcop](#), is how the parameter range of Θ (the para[1] variable) is controlled—here the parameter remains untransformed but the lower domain edge is truncated

by the return of infinity (return(Inf)). The getstat argument is only for short circuiting the objective so that objfunc can be used to compute the three statistics after the optimal parameters are found.

```
"objfunc" <- function(para, stat=NA, as.sample=FALSE, getstat=FALSE, ...) {
  if(as.sample) {
    return(c( giniCOP(para=para, as.sample=TRUE),
              nuskewCOP(para=para, as.sample=TRUE),
              nustarCOP(para=para, as.sample=TRUE)))
  }
  para[1] <- ifelse(para[1] < 0, return(Inf), para[1]) # edge check
  para[2:3] <- pnorm(para[2:3]) # detransform
  para <- list(cop1=HRCop, para1=para[1], alpha=para[2], beta=para[3])
  hp <- c( giniCOP(composite1COP, para),
           nuskewCOP(composite1COP, para),
           nustarCOP(composite1COP, para))
  if(getstat) return(hp) # short circuit to get the statistics out
  dv <- stat; dv[dv == 0] <- 1 # changing dv "adapts" the error to
  return(sqrt(sum(((stat-hp)/dv)^2))) # trap division by zero
}
```

The parameter estimation proceeds in the following code. The sample statistics (or target.stats) are computed and subsequently passed to the optimization using the popstat argument. Notice also the use of the qnorm() function in R to transform the initial guess $\alpha = \beta = 1/2$ into a domain more easily handled by the optimizer (optim() function in R). The transformed optimal parameters are computed, and then the values of the three statistics for the fit are computed. Lastly, a **copBasic** parameter object fit para is created, which can be used for later comparisons.

```
txt <- c("GiniGamma", "NuSkew", "NuStar")
target.stats <- objfunc(sampleUV, as.sample=TRUE); names(target.stats) <- txt
raw.fit.para <- copBasic.fitpara.beta(popstat=target.stats, statf=objfunc,
  par.init=c(1, qnorm(0.5), qnorm(0.5)), cop=composite1COP, paradim=3)
fit.stats <- objfunc(raw.fit.para, getstat=TRUE); names(fit.stats) <- txt
fit.para <- list(cop1=HRCop, para1=raw.fit.para[1],
  alpha=pnorm(raw.fit.para[2]), beta=pnorm(raw.fit.para[3]))
```

The optimization is completed. It is informative to see what the simulation of the fitted copula looks like. Note: this particular example suffers from identifiability problems between the parameters—meaning that local minima exist or that satisfactory solutions using different parameters than used to generate the random sample can occur. The same seed is used so that one-to-one comparison of points can visually be made with the [simCOP](#) function call.

```
set.seed(1999) # This value will be used again (see below)
sampleUV <- simCOP(n=n, cop=composite1COP, para=fit.para,
  ploton=FALSE, pch=16, cex=0.5, col=2, snv=TRUE) # red dots
```

The visual comparison is qualitative. The tabular comparison of the sample statistics to those of the fitted model shows that perhaps an acceptable local minima has been attained in terms of “fit” but the subsequent comparison of the parameters of the population used to generate the sample

and those of the fitted model seemingly depart substantially in the $\alpha \rightarrow 0$ parameter of the copula composition. The tail dependency parameters are similar, but further goodness-of-fit check is not made.

```
#      GiniGamma      NuSkew      NuStar
print(target.stats) #      0.5219027      -0.1940361      0.6108319
print(fit.stats)    #      0.5182858      -0.1938848      0.6159566

# Parameter Alpha      Beta
print(ls.str(pop.para)) #      4.00      0.14      0.350 # given
print(ls.str(fit.para)) #      11.2      0.187      0.427 # one solution

# Tail Dependency Parameters
taildepCOP(cop=composite1COP, para=pop.para) # lower=0 : upper=0.5838
taildepCOP(cop=composite1COP, para=fit.para) # lower=0 : upper=0.5714(est.)
```

The demonstration ends with the comparison of the two asymmetrical density contours.

```
densityCOPplot(cop=composite1COP, para=pop.para, contour.col=3)
densityCOPplot(cop=composite1COP, para=fit.para, contour.col=2,
               ploton=FALSE)
```

Author(s)

W.H. Asquith

Examples

```
# See the Note section
```

COPinv

The Inverse of a Copula for V with respect to U

Description

Compute the *inverse of a copula* for V with respect to U given t or

$$t = \mathbf{C}(u=U, v) \rightarrow \mathbf{C}^{(-1)}(u=U, t) = v,$$

and solving for v . Nelsen (2006, p. 12) does not so name this function as an “inverse.” The COPinv function is internally used by [level.curvesCOP](#) and [level.curvesCOP2](#). A common misapplication that will puzzle the user (including the developer after long breaks from package use) is that the following call and error message are often seen, if `silent=FALSE`:

```
COPinv(0.2, 0.25, cop=PSP)
# Error in uniroot(func, interval = c(lo, 1), u = u, LHS = t, cop = cop, :
# f() values at end points not of opposite sign
# [1] NA
```

This is a harmless error in the sense that COPinv is functioning properly. One can not invert a copula for $u < t$ and for $u = t$ the $v = 1$ because of fundamental copula properties.

Usage

```
COPinv(cop=NULL, u, t, para=NULL, silent=TRUE, ...)
```

Arguments

<code>cop</code>	A copula function;
<code>u</code>	Nonexceedance probability u in the X direction;
<code>t</code>	Nonexceedance probability level t ;
<code>para</code>	Vector of parameters or other data structures, if needed, to pass to the copula;
<code>silent</code>	The argument of the same name given over to <code>try()</code> wrapping the <code>uniroot()</code> operation; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for v are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[COP](#), [COPinv2](#), [level.curvesCOP](#), [level.curvesCOP2](#)

Examples

```
COPinv(cop=N4212cop, para=2, u=0.5, t=0.2)
```

COPinv2

The Inverse of a Copula for U with respect to V

Description

Compute the *inverse of a copula* for U with respect to V given t or

$$t = \mathbf{C}(u, v=V) \rightarrow \mathbf{C}^{(-1)}(v=V, t) = u,$$

and solving for u . Nelsen (2006, p. 12) does not so name this function as an “inverse.” The `COPinv2` function is internally used by [level.curvesCOP2](#). A common misapplication that will puzzle the user (including the developer after long breaks from package use) is that the following call and error message are often seen, if `silent=FALSE`:

```

COPinv2(0.2, 0.25, cop=PSP)
# Error in uniroot(func, interval = c(lo, 1), u = u, LHS = t, cop = cop, :
# f() values at end points not of opposite sign
# [1] NA

```

This is a harmless error in the sense that COPinv2 is functioning properly. One can not invert a copula for $v < t$ and for $v = t$ the $u = 1$ because of fundamental copula properties.

Usage

```
COPinv2(cop=NULL, v, t, para=NULL, silent=TRUE, ...)
```

Arguments

cop	A copula function;
v	Nonexceedance probability v in the Y direction;
t	Nonexceedance probability in t ;
para	Vector of parameters or other data structures, if needed, to pass to the copula;
silent	The argument of the same name given over to try() wrapping the uniroot() operation; and
...	Additional arguments to pass to the copula.

Value

Value(s) for u are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[COP](#), [COPinv](#), [level.curvesCOP](#), [level.curvesCOP2](#)

Examples

```
# See those for COPinv as they are the same by analogy.
```

densityCOP

*Density of a Copula***Description**

Numerically estimate the *copula density* for a sequence of u and v probabilities for which each sequence has equal steps that are equal to $\Delta(uv)$. The density $c(u, v)$ of a copula $\mathbf{C}(u, v)$ is numerically estimated by

$$c(u, v) = [\mathbf{C}(u_2, v_2) - \mathbf{C}(u_2, v_1) - \mathbf{C}(u_1, v_2) + \mathbf{C}(u_1, v_1)] / [\Delta(uv) \times \Delta(uv)],$$

where $c(u, v) \geq 0$ (see Nelsen, 2006, p. 10; [densityCOPplot](#)). The *joint density* can be defined by the coupla density for continuous variables and is the ratio of the joint density function $f(x, y)$ for random variables X and Y to the product of the marginal densities ($f_x(x)$ and $f_y(y)$):

$$c(F_x(x), F_y(y)) = \frac{f(x, y)}{f_x(x)f_y(y)},$$

where $F_x(x)$ and $F_y(y)$ are the respective cumulative distribution functions of X and Y , and lastly $u = F_x(x)$ and $v = F_y(y)$.

Usage

```
densityCOP(u,v, cop=NULL, para=NULL, deluv=.Machine$double.eps^0.25,
           truncate.at.zero=TRUE, the.zero=0, sumlogs=FALSE, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>deluv</code>	The change in the sequences $\{u, v\} \mapsto \delta, \dots, 1 - \delta$; $\delta = \Delta(uv)$ probabilities;
<code>truncate.at.zero</code>	A density must be $c(u, v) \geq 0$, but because this computation is based on a rectangular approximation and not analytical, there exists a possibility that very small rectangles could result in numerical values in $\mathbb{R}$ that are less than zero. This usually can be blamed on rounding. This logical if TRUE truncates computed densities to zero, and the default assumes that the user is providing a proper copula. A FALSE value is used by the function isfuncCOP ;
<code>the.zero</code>	The value for “the zero” where a small number might be useful for pseudo-maximum likelihood estimation using <code>sumlogs</code> ;
<code>sumlogs</code>	Return the $\sum \log c(u, v; \Theta)$ where Θ are the parameters in <code>para</code> and this feature is provided for mleCOP ; and
<code>...</code>	Additional arguments to pass to the copula function.

Value

Value(s) for $c(u, v)$ are returned.

Note

The **copBasic** package does not currently have copula densities as analytical solutions implemented. This is because initial design decisions were entirely on cumulative distribution function (CDF) representations of the copula.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[simCOP](#), [densityCOPplot](#), [kullCOP](#), [mleCOP](#)

Examples

```
## Not run:
# Joe (2014, p. 164) shows the closed form copula density for the Plackett.
"dPLACKETTcop" <- function(u,v,para) {
  uv <- u*v; upv <- u + v; eta <- para - 1
  A <- para*(1+eta*(upv - 2*uv)); B <- ((1+eta*upv)^2 - 4*para*eta*uv)^(3/2)
  return(A/B)
}
dPLACKETTcop(0.32, 0.74, para=1.3) # 0.9557124
densityCOP( 0.32, 0.74, cop=PLcop, para=1.3) # 0.9557153
## End(Not run)

## Not run:
# Joe (2014, p. 165) shows the corner densities of the Plackett as Theta.
# copBasic uses numerical density estimation and not analytical formula.
eps <- .Machine$double.eps
densityCOP(0,0, cop=PLcop, para=4) # 3.997073 (default eps^0.25)
densityCOP(1,1, cop=PLcop, para=4) # 3.997073 (default eps^0.25)
densityCOP(1,1, cop=PLcop, para=4, deluv=eps) # 0 (silent failure)
densityCOP(1,1, cop=PLcop, para=4, deluv=eps^0.5) # 4.5
densityCOP(1,1, cop=PLcop, para=4, deluv=eps^0.4) # 4.002069
densityCOP(1,1, cop=PLcop, para=4, deluv=eps^0.3) # 3.999513
# So, we see that the small slicing does have an effect, the default of 0.25 is
# intended for general application by being away enough from machine limits.
## End(Not run)

## Not run:
# Joe (2014, p. 170) shows a closed form copula density for "Bivariate Joe/B5" copula
```

```

"dJOEB5cop" <- function(u, v, para) {
  up <- (1-u)^para; vp <- (1-v)^para; eta <- para - 1
  A <- (up + vp - up*vp); B <- (1-u)^eta * (1-v)^eta; C <- (eta + A)
  return(A^(-2 + 1/para) * B * C)
}
densityCOP(0.32, 0.74, cop=JOcopB5, para=1.3) # 0.9410653
dJOEB5cop( 0.32, 0.74, para=1.3)             # 0.9410973
## End(Not run)

```

densityCOPplot

Contour Density Plot of a Copula

Description

Generate a *contour density plot* after the advocacy of Joe (2014, pp. 9–15). Such graphics are plots of *scaled copula densities* ($c^*(u, v)$, bivariate herein) that are copula densities scaled to the standard normal distribution $\sim N(0,1)$ margins. Joe (2014) repeatedly emphasizes such plots in contrast to the uniform distribution $\sim U(0,1)$ margins. Nelsen (2006) does not discuss such scaling but seemingly Nelsen's objectives for his book were somewhat different.

The density of copula $C(u, v)$ is numerically estimated by

$$c(u, v) = [C(u_2, v_2) - C(u_2, v_1) - C(u_1, v_2) + C(u_1, v_1)] / [\Delta(uv) \times \Delta(uv)],$$

where $c(u, v) \geq 0$ (see Nelsen, 2006, p. 10; [densityCOP](#)). Given a numerically estimated quantity $c^*(u, v) = c(u, v) \times \phi(\Phi^{-1}(u)) \times \phi(\Phi^{-1}(v))$ for copula density $c(u, v)$, a grid of the $c^*(u, v)$ values can be contoured by the `contour()` function in R. The density function of the $N(0,1)$ is $\phi(z)$ for standard normal variate z and the quantile function of the $N(0,1)$ is $\Phi^{-1}(t)$ for nonexceedance probability t .

A grid (matrix) of $c(u, v)$ or $c^*(u, v)$ is defined for sequence of u and v probabilities for which each sequence has equal steps that are equal to $\Delta(uv)$. This function has as focus on plotting of the contour lines of $c^*(u, v)$ but the R matrix of either $c(u, v)$ or $c^*(u, v)$ can be requested on return. For either matrix, the `colnames()` and `rownames()` (the R functions) are set equal to the sequence of u and v , respectively. Neither the column or row names are set to the standard normal variates for the matrix of $c^*(u, v)$, the names remain in terms of nonexceedance probability.

For plotting and other uses of normal scores of data, Joe (2014, p. 245) advocates that one should use the plotting position formula $u_i = (i - 1/2)/n$ (*Hazen plotting position*) for normal scores $z_i = \Phi^{-1}(u_i)$ in preference to $i/(n + 1)$ (*Weibull plotting position*) because $n^{-1} \sum_{i=1}^n z_i^2$ is closer to unity. Other examples of Joe's advocacy for the Hazen plotting positions are available (Joe, 2014, pp. 9, 17, 245, 247–248).

Usage

```

densityCOPplot(cop=NULL, para=NULL, deluv=0.002,
  getmatrix=c("none", "cdenzz", "cden"), n=0,
  ploton=TRUE, snv=TRUE, origins=TRUE,
  contour.col=1, contour.lwd=1.5, ...)

```

Arguments

<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>deluv</code>	The change in the sequences $\{u, v\} \mapsto \delta, \dots, 1 - \delta$; $\delta = \Delta(uv)$ probabilities;
<code>getmatrix</code>	A trigger on whether the density matrix is to be returned. The option <code>cdenzz</code> returns the density scaled by the two standard normal densities ($c^*(u, v)$), whereas the option <code>cden</code> returns simply the copula density ($c(u, v)$);
<code>ploton</code>	A logical to toggle on the plot;
<code>snv</code>	A logical to toggle standard normal variates for the axes;
<code>origins</code>	A logical to plot the origin lines, if and only if <code>snv</code> is true;
<code>contour.col</code>	The color of the contour lines, which corresponds to the <code>col</code> argument of the contour function in R;
<code>contour.lwd</code>	The width of the contour lines, which corresponds to the <code>lwd</code> argument of the contour function in R;
<code>n</code>	An optional sample size for which simulation of this many values from the copula will be made by simCOP and drawn; and
<code>...</code>	Additional arguments to pass to the copula function and to the contour function in R (e.g. to turn off labeling of contours add <code>drawlabels=FALSE</code>).

Value

This is a high-level function used for its side effects; an R matrix can be triggered, however, as a returned value.

Note

Joe (2014, p. 244) says “if [density] plots show multimodality, then multivariate techniques of mixture models, cluster analysis[,] and nonparametric functional data analysis might be more appropriate” than relatively straightforward parametric copula models.

Author(s)

W.H. Asquith

References

- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
 Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[simCOP](#), [densityCOP](#)

Examples

```
## Not run:
# Joe (2014, p. 5) names rMTCJ = reflected Mardia-Takahasi-Cook-Johnson copula
"rMTCJ" <- function(u, v, para, ...) {
  u + v - 1 + ((1-u)^(-para) + (1-v)^(-para) - 1)^(-1/para)
} # Survival Copula ("reflected" in Joe's terms)
densityCOPplot(cop=rMTCJ, para=1.0760, n=9000, snv=TRUE)
# Density plot matches that shown by Joe (2014, p. 11, fig. 1.2, lower left plot)
# for a Spearman Rho equaling 0.5. Let us compute then Rho:
rhoCOP(cop=rMTCJ, para=1.076075) # 0.4999958

# Now, let us get really wild with a composition with TWO modes!
# This example also proves that the grid orientation is consistent with respect
# to the orientation of the simulations.
para <- list(alpha=0.15, beta=0.90, kappa=0.06, gamma=0.96,
             cop1=GHCop, cop2=PLACKETTcop, para1=5.5, para2=0.07)
densityCOPplot(cop=composite2COP, para=para, n=9000)

# Now, let us hack back to a contour density plot with U(0,1) and not N(0,1) margins
# just so show that capability exists, but emphasis of densityCOPplot is clearly
# on Joe's advocacy, because it does not have a default trigger to use U(0,1) margins.
set.seed(12)
H <- densityCOPplot(cop=PLACKETTcop, para=41.25, getmatrix="cdenz", n=1000, snv=FALSE)
set.seed(12)
UV <- simCOP(cop=PLACKETTcop, para=41.25, n=1000, col=8, snv=FALSE)
U <- as.numeric(colnames(H)); V <- as.numeric(rownames(H))
contour(x=U, y=V, z=t(H), lwd=1.5, cex=2, add=TRUE, col=2) #
## End(Not run)

## Not run:
set.seed(12)
UV <- rCOP(400, cop=PSP, pch=16, col=8, n=400)
CL <- mleCOP(UV, cop=CLcop, interval=c(1, 20))
JO <- mleCOP(UV, cop=JOcopB5, interval=c(0.1, 20))
PL <- mleCOP(UV, cop=PLcop, interval=c(0.1, 20))

AICs <- c(CL$AIC, JO$AIC, PL$AIC)
names(AICs) <- c("Clayton", "Joe(B5)", "Plackett")
print(round(AICs, digits=2))
# Clayton   Joe(B5)   Plackett
# -156.77    -36.91   -118.38
# So, we conclude Clayton is the best fit of the three.

plot(qnorm(UV[,1]), qnorm(UV[,2]), pch=16, col=8, cex=0.5,
     xlab="Standard normal variate of U", xlim=c(-3, 3),
     ylab="Standard normal variate of V", ylim=c(-3, 3))
densityCOPplot(cop=PSP, contour.col=grey(0.5), lty=2,
               contour.lwd=3.5, ploton=FALSE, snv=TRUE)
densityCOPplot(cop=CLcop, para=CL$para,
               contour.col=2, ploton=FALSE, snv=TRUE)
densityCOPplot(cop=JOcopB5, para=JO$para,
               contour.col=3, ploton=FALSE, snv=TRUE)
```

```

densityCOPplot(cop=PLcop,      para=PL$para,
               contour.col=4,   ploton=FALSE, snv=TRUE) #
## End(Not run)

```

derCOP

Numerical Derivative of a Copula for V with respect to U

Description

Compute the numerical partial derivative of a copula, which is a *conditional distribution function*, according to Nelsen (2006, pp. 13, 40–41) with respect to u :

$$0 \leq \frac{\delta}{\delta u} \mathbf{C}(u, v) \leq 1,$$

or

$$\Pr[V \leq v \mid U = u] = \mathbf{C}_{2|1}(v \mid u) = \lim_{\Delta u \rightarrow 0} \frac{\mathbf{C}(u + \Delta u, v) - \mathbf{C}(u, v)}{\Delta u},$$

which is to read as the probability that $V \leq v$ given that $U = u$ and corresponds to the `derdir="left"` mode of the function. For `derdir="right"`, we have

$$\Pr[V \leq v \mid U = u] = \lim_{\Delta u \rightarrow 0} \frac{\mathbf{C}(u, v) - \mathbf{C}(u - \Delta u, v)}{\Delta u},$$

and for `derdir="center"` (the usual method of computing a derivative), the following results

$$\Pr[V \leq v \mid U = u] = \lim_{\Delta u \rightarrow 0} \frac{\mathbf{C}(u + \Delta u, v) - \mathbf{C}(u - \Delta u, v)}{2\Delta u}.$$

The “with respect to V ” versions are available under [derCOP2](#).

Copula derivatives ($\delta \mathbf{C} / \delta u$ or say $\delta \mathbf{C} / \delta v$ [derCOP2](#)) are non-decreasing functions meaning that if $v_1 \leq v_2$, then $\mathbf{C}(u, v_2) - \mathbf{C}(u, v_1)$ is a non-decreasing function in u , thus

$$\frac{\delta(\mathbf{C}(u, v_2) - \mathbf{C}(u, v_1))}{\delta u}$$

is non-negative, which means

$$\frac{\delta \mathbf{C}(u, v_2)}{\delta u} \geq \frac{\delta \mathbf{C}(u, v_1)}{\delta u} \text{ for } v_2 \geq v_1.$$

Usage

```

derCOP(cop=NULL, u, v, delu=.Machine$double.eps^0.50,
       derdir=c("left", "right", "center"), ...)

```

Arguments

<code>cop</code>	A copula function;
<code>u</code>	Nonexceedance probability u in the X direction. If the length of <code>u</code> is unity, then the length of <code>v</code> can be arbitrarily long. If the length of <code>u</code> is not unity, then the length of <code>v</code> should be the same, and if not, then only the first value in <code>v</code> is silently used;
<code>v</code>	Nonexceedance probability v in the Y direction (see previous comment on <code>u</code>);
<code>delu</code>	The Δu interval for the derivative;
<code>derdir</code>	The direction of the derivative as described above. Default is <code>left</code> but internally any setting can be temporarily suspended to avoid improper computations (see source code); and
<code>...</code>	Additional arguments to pass such as the parameters often described in <code>para</code> arguments of other copula functions. (The lack of <code>para=NULL</code> for <code>derCOP</code> and <code>derCOP2</code> was either design oversight or design foresight but regardless it is too late to enforce package consistency in this matter.)

Value

Value(s) for the partial derivative are returned.

Note

A known caveat of the current implementation of the copula derivative is that there is a chance that the Δu will span a singularity or discontinuous (or nearly so) portion of a copula should it have a property of singularity (or nearly so). The `delu` is chosen small so the chance is mitigated to be a small change and certainly appear to work throughout the examples herein. It is not decided whether a derivative from the positive side (`dir="left"`), when failing should switch over to a computation from the negative side (`dir="right"`). The distinction is important for the computation of the inverse of the derivative `derCOPinv` because the solution finder needs a sign reversal to work.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

`derCOPinv`, `derCOP2`

Examples

```
derCOP(cop=W, 0.4, 0.6); derCOP(cop=P, 0.4, 0.6); derCOP(cop=M, 0.4, 0.6)

lft <- derCOP(cop=PSP, 0.4, 0.6, derdir="left" )
rgt <- derCOP(cop=PSP, 0.4, 0.6, derdir="right" )
```

```

cnt <- derCOP(cop=PSP, 0.4, 0.6, derdir="center")
cat(c(lft,rgt,cnt,"\n"))
#stopifnot(all.equal(lft,rgt), all.equal(lft,cnt))

# Let us contrive a singularity through this NOT A COPULA in the function "afunc".
"afunc" <- function(u,v, ...) return(ifelse(u <= 0.5, sqrt(u^2+v^2), P(u,v,...)))
lft <- derCOP(cop=afunc, 0.5, 0.67, derdir="left" )
rgt <- derCOP(cop=afunc, 0.5, 0.67, derdir="right" )
cnt <- derCOP(cop=afunc, 0.5, 0.67, derdir="center")
cat(c(lft,rgt,cnt,"\n")) # The "right" version is correct.

```

derCOP2

Numerical Derivative of a Copula for U with respect to V

Description

Compute the numerical partial derivative of a copula, which is a *conditional distribution function*, according to Nelsen (2006, pp. 13, 40–41) with respect to v :

$$0 \leq \frac{\delta}{\delta v} \mathbf{C}(u, v) \leq 1,$$

or

$$\Pr[U \leq u \mid V = v] = \mathbf{C}_{1|2}(u \mid v) = \lim_{\Delta v \rightarrow 0} \frac{\mathbf{C}(u, v + \Delta v) - \mathbf{C}(u, v)}{\Delta v},$$

which is to read as the probability that $U \leq u$ given that $V = v$ and corresponds to the `derdir="left"` mode of the function. For `derdir="right"`, the following results

$$\Pr[U \leq u \mid V = v] = \lim_{\Delta v \rightarrow 0} \frac{\mathbf{C}(u, v) - \mathbf{C}(u, v - \Delta v)}{\Delta v},$$

and for `derdir="center"` (the usual method of computing a derivative), the following results

$$\Pr[U \leq u \mid V = v] = \lim_{\Delta v \rightarrow 0} \frac{\mathbf{C}(u, v + \Delta v) - \mathbf{C}(u, v - \Delta v)}{2\Delta v}.$$

The “with respect to U ” versions are under [derCOP](#).

Copula derivatives ($\delta \mathbf{C} / \delta v$ or say $\delta \mathbf{C} / \delta u$ [derCOP](#)) are non-decreasing functions meaning that if $u_1 \leq u_2$, then $\mathbf{C}(u_2, v) - \mathbf{C}(u_1, v)$ is a non-decreasing function in v , thus

$$\frac{\delta (\mathbf{C}(u_2, v) - \mathbf{C}(u_1, v))}{\delta v}$$

is non-negative, which means

$$\frac{\delta \mathbf{C}(u_2, v)}{\delta v} \geq \frac{\delta \mathbf{C}(u_1, v)}{\delta v} \text{ for } u_2 \geq u_1.$$

Usage

```
derCOP2(cop=NULL, u, v, delv=.Machine$double.eps^0.50,
        derdir=c("left", "right", "center"), ...)
```

Arguments

<code>cop</code>	A copula function;
<code>u</code>	Nonexceedance probability u in the X direction. If the length of <code>u</code> is unity, then the length of <code>v</code> can be arbitrarily long. If the length of <code>u</code> is not unity, then the length of <code>v</code> should be the same and if not only the first value in <code>v</code> will be silently used;
<code>v</code>	Nonexceedance probability v in the Y direction (see previous comment on <code>u</code>);
<code>delv</code>	The Δv interval for the derivative;
<code>derdir</code>	The direction of the derivative as described above. Default is <code>left</code> but internally any setting can be temporarily suspended to avoid improper computations (see source code); and
<code>...</code>	Additional arguments to pass such as the parameters often described in <code>para</code> arguments of other copula functions. (The lack of <code>para=NULL</code> for derCOP and derCOP2 was either design oversight or design foresight but regardless it is too late to enforce package consistency in this matter.)

Value

Value(s) for the partial derivative are returned.

Note

A known caveat of the current implementation of the copula derivative is that there is a chance that the Δv will span a singularity or discontinuous (or nearly so) portion of a copula should it have a property of singularity (or nearly so). The `delv` is chosen small so the chance is mitigated to be a small change and certainly seems to work throughout the examples herein. It is not decided whether a derivative from the positive side (`dir="left"`), when failing should switch over to a computation from the negative side (`dir="right"`). The distinction is important for the computation of the inverse of the derivative [derCOPinv2](#) because the solution finder needs a sign reversal to work.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[derCOPinv2](#), [derCOP](#)

Examples

```

derCOP2(cop=W, 0.4, 0.6); derCOP2(cop=P, 0.4, 0.6); derCOP2(cop=M, 0.4, 0.6)

lft <- derCOP2(cop=P, 0.4, 0.6, derdir="left" )
rgt <- derCOP2(cop=P, 0.4, 0.6, derdir="right" )
cnt <- derCOP2(cop=P, 0.4, 0.6, derdir="center")
cat(c(lft, rgt, cnt, "\n"))
# stopifnot(all.equal(lft, rgt), all.equal(lft, cnt))

# Let us contrive a singularity though this NOT A COPULA in the function "afunc".
"afunc" <- function(u,v, ...) return(ifelse(u <= 0.5, sqrt(u^2+v^2), P(u,v,...)))
lft <- derCOP2(cop=afunc, 0.67, 0.5, derdir="left" )
rgt <- derCOP2(cop=afunc, 0.67, 0.5, derdir="right" )
cnt <- derCOP2(cop=afunc, 0.67, 0.5, derdir="center")
cat(c(lft,rgt,cnt,"\n")) # For this example, all are correct (see derCOP examples)

```

derCOPinv

Numerical Derivative Inverse of a Copula for V with respect to U

Description

Compute the inverse of a numerical partial derivative for V with respect to U of a copula, which is a *conditional quantile function* for nonexceedance probability t , or

$$t = c_u(v) = \mathbf{C}_{2|1}^{(-1)}(v \mid u) = \frac{\delta \mathbf{C}(u, v)}{\delta u},$$

and solving for v . Nelsen (2006, pp. 13, 40–41) shows that this inverse is quite important for random variable generation using the *conditional distribution method*. This function is not vectorized and will not be so.

Usage

```

derCOPinv(cop=NULL, u, t, trace=FALSE,
          delu=.Machine$double.eps^0.50, para=NULL, ...)

```

Arguments

cop	A copula function;
u	A single nonexceedance probability u in the X direction;
t	A single nonexceedance probability level t ;
trace	A logical controlling a message on whether the signs on the uniroot are the same—this is helpful in exploring the numerical derivative limits of a given implementation of a copula.
delu	The Δu interval for the derivative;
para	Vector of parameters or other data structures, if needed, to pass to cop; and
...	Additional arguments to pass to the copula.

Value

Value(s) for the derivative inverse are returned.

Note

An Educational Opportunity—The *Farlie–Gumbel–Morgenstern* copula $\mathbf{FGM}(u, v)$ ([FGMcop](#)) (Joe, 2014, p. 213) is

$$\mathbf{FGM}(u, v; \Theta) = uv[1 + \Theta(1 - u)(1 - v)],$$

where $-1 \leq \Theta \leq 1$ has analytical solutions to the *conditional cumulative distribution function* (CCDF) $\mathbf{C}_{2|1}(v | u)$ as

$$\mathbf{C}_{2|1}(v | u) = v[1 + \Theta(1 - v)(1 - 2u)],$$

and the inverse of the CCDF as

$$\mathbf{C}_{2|1}(v | u) = \frac{[1 + \Theta(1 - 2u)] - \sqrt{[1 + \Theta(1 - 2u)]^2 - 4t(1 - 2u)}}{2\Theta(1 - 2u)}.$$

These three functions for the copula can be defined in R by

```
"FGMcop"      <- function(u,v, para=NULL, ...) u*v*(1 + para*(1-u)*(1-v) )
"joeFGMder"   <- function(u,v, para=NULL, ...)  v*(1 + para*(1-v)*(1-2*u))
"joeFGMderinv" <- function(u,t, para=NULL, ...) { K <- (1-2*u)
  ((1 + para*K) - sqrt((1 + para*K)^2 - 4*t*K))/(2*para*K)
}
```

The $\mathbf{C}_{2|1}^{(-1)}(v | u)$ is critical for simulation by the conditional simulation method. Although exclusively for simulation, **copBasic** uses inversion of the numerical derivative, the **FGM** copula has three representations of supposedly the same analytical algorithm for simulation in the literature (Durante, 2007; Johnson, 1987; Nelsen, 2006). An opportunity for comparison is thus available.

The three analytical algorithms for nonexceedance probability t given u by mathematics and code, following Durante (2007, p. 245), are

$$\begin{aligned} A &= \Theta(1 - 2u) - 1, \\ B &= \sqrt{A^2 - 4t(A + 1)}, \text{ and} \\ v &= 2t/(B - A), \end{aligned}$$

and in R, this “Durante algorithm” is

```
"durFGMderinv" <- function(u,t, para=NULL, ...) { # Durante (2007, p. 245)
  A <- para*(1-2*u) - 1; B <- sqrt(A^2 - 4*t*(A+1)); return(2*t/(B - A))
}
```

and, letting $K = (2u - 1)$, following Johnson (1987, p. 185)

$$\begin{aligned} A &= K\Theta - 1 \\ B &= \sqrt{1 - 2K\Theta + (K\Theta)^2 + 4tK\Theta} \\ v &= 2t/(B - A) \end{aligned}$$

and in R, this “Johnson algorithm” is

```
"jonFGMderinv" <- function(u,t, para=NULL, ...) { # Johnson (1987, p. 185)
  K <- (2*u - 1)
  A <- K*para - 1; B <- sqrt(1 - 2*K*para + (K*para)^2 + 4*t*K*para)
  2*t/(B - A)
}
```

and finally following Nelsen (2006, p. 87)

$$A = 1 + \theta(1 - 2u),$$

$$B = \sqrt{A^2 - 4t(A - 1)}, \text{ and}$$

$$v = 2t/(B + A),$$

and in R, this “Nelsen algorithm” is

```
"nelFGMderinv" <- function(u,t, para=NULL, ...) { # Nelsen (2006, p. 87)
  A <- 1 + para*(1-2*u); B <- sqrt(A^2 - 4*t*(A-1)); return(2*t/(B + A))
}
```

With appropriate code now available, two comparisons can be made in the following section.

Conditional Cumulative Distribution Function—A comparison of the analytical **FGM**_(u,v) derivative shows that Joe’s equation is congruent with the numerical derivative of **copBasic**:

```
joeFGMder(0.8, 0.44, para=0.78) # 0.3246848 (Joe, 2014)
derCOP( 0.8, 0.44, para=0.78, cop=FGMcop) # 0.3246848 (copBasic )
```

and the result will be used in the computations that follow.

A comparison for $t = 0.3246848$ of the analytical inverse and the numerical optimization of the numerical derivative of **copBasic** is

```
joeFGMderinv(0.8, 0.3246848, para=0.78) # 0.5327603
derCOPinv( 0.8, 0.3246848, para=0.78, cop=FGMcop) # 0.4399934 --> 0.44
```

where obviously, the two results are not in agreement—so something is amiss. Because many examples in this documentation clearly demonstrate numerical reliability, a tentative conclusion is that Joe’s listed equation must be in error. Let us check this hypothesis against the three other sources:

```
durFGMderinv(0.8, 0.3246848, para=0.78) # 0.2074546 (Durante, 2007)
jonFGMderinv(0.8, 0.3246848, para=0.78) # 0.44 (Johnson, 1987)
nelFGMderinv(0.8, 0.3246848, para=0.78) # 0.44 (Nelsen, 2006)
```

The result from Durante (2007) is different from both Joe (2014) and from **copBasic**. However, the Johnson (1987) and Nelsen (2006) versions are equivalent and congruent to **copBasic** with the *distinctly different* numerical methods of derCOPinv. These incongruent results demonstrate that care is needed when navigating the copula literature and the usefulness of the **copBasic**-style implementation of copula theory. In words, these computations show that the $t \approx 32$ nd percentile of the **FGM** copula given that the 80th percentile in U is about the 44th percentile of V .

Author(s)

W.H. Asquith

References

- Durante, F., 2007, Families of copulas, Appendix C, *in* Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, *Extremes in Nature—An approach using copulas*: Springer, 289 p.
- Joe, H., 2014, *Dependence modeling with copulas*: Boca Raton, CRC Press, 462 p.
- Johnson, M.E., 1987, *Multivariate statistical simulation*: New York, John Wiley, 230 p.
- Nelsen, R.B., 2006, *An introduction to copulas*: New York, Springer, 269 p.
- Zhang, L., and Singh, V.P., 2019, *Copulas and their applications in water resources engineering*: Cambridge University Press, ISBN 978–1–108–47425–2.

See Also

[derCOP](#)

Examples

```
u <- runif(1); t <- runif(1)
derCOPinv(u,t, cop=W ) # perfect negative dependence
derCOPinv(u,t, cop=P ) # independence
derCOPinv(u,t, cop=M ) # perfect positive dependence
derCOPinv(u,t, cop=PSP) # a parameterless copula example
## Not run:
# Simulate 500 values from product (independent) copula
plot(NA,NA, type="n", xlim=c(0,1), ylim=c(0,1), xlab="U", ylab="V")
for(i in 1:500) {
  u <- runif(1); t <- runif(1)
  points(u, derCOPinv(cop=P, u, t), cex=0.5, pch=16) # black dots
}
# Now simulate 500 from the Nelsen 4.2.12 copula.
for(i in 1:500) {
  u <- runif(1); t <- runif(1)
  points(u,derCOPinv(cop=N4212cop,para=9.3,u,t), cex=2, pch=16, col=2) # red dots
} #
## End(Not run)

## Not run:
# Zhang and Singh (2019) exam. 3.23, p. 105 show the application of the
# derivative inversion C2|1 for u=0.6036 and t=0.6036 ---> v = 0.4719
derCOPinv( cop=CLcop, 0.6036, 0.4028, para=0.5) # 0.4719 for C2|1
derCOPinv2(cop=CLcop, 0.6036, 0.4028, para=0.5) # 0.4719 for C1|2
# and C2|1 and C1|2 are equal because the copula has permutation symmetry
isCOP.permsym(cop=CLcop, para=0.5) # TRUE
## End(Not run)
```

derCOPinv2

*Numerical Derivative Inverse of a Copula for U with respect to V***Description**

Compute the inverse of a numerical partial derivative for U with respect to V of a copula, which is a *conditional quantile function* for nonexceedance probability t , or

$$t = c_v(u) = \mathbf{C}_{1|2}^{(-1)}(u | v) = \frac{\delta \mathbf{C}(u, v)}{\delta v},$$

and solving for u . Nelsen (2006, pp. 13, 40–41) shows that this inverse is quite important for random variable generation using the *conditional distribution method*. This function is not vectorized and will not be so.

Usage

```
derCOPinv2(cop=NULL, v, t, trace=FALSE,
           delv=.Machine$double.eps^0.50, para=NULL, ...)
```

Arguments

cop	A copula function;
v	A single nonexceedance probability v in the Y direction;
t	A single nonexceedance probability level t ;
trace	A logical controlling a message on whether the signs on the uniroot are the same—this is helpful in exploring the numerical derivative limits of a given implementation of a copula.
delv	The Δv interval for the derivative;
para	Vector of parameters or other data structure, if needed, to pass to cop; and
...	Additional arguments to pass to the copula.

Value

Value(s) for the derivative inverse are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[derCOP2](#)

Examples

```

u <- runif(1); t <- runif(1)
derCOPinv2(u,t, cop=W) # perfect negative dependence
derCOPinv2(u,t, cop=P) # independence
derCOPinv2(u,t, cop=M) # perfect positive dependence
derCOPinv2(u,t, cop=PSP) # a parameterless copula example
## Not run:
# Simulate 500 values from product (independent) copula
plot(NA,NA, type="n", xlim=c(0,1), ylim=c(0,1), xlab="U", ylab="V")
for(i in 1:500) {
  v <- runif(1); t <- runif(1)
  points(derCOPinv2(cop=P, v, t),v, cex=0.5, pch=16) # black dots
}
# Simulate 500 of a Frechet Family copula and note crossing singularities.
for(i in 1:500) {
  v <- runif(1); t <- runif(1)
  u <- derCOPinv2(v, t, cop=FRECHETcop, para=list(alpha=0.7, beta=0.169))
  points(u,v, cex=2, pch=16, col=2) # red dots
} #
## End(Not run)

```

diagCOP

The Diagonals of a Copula

Description

Compute the *primary diagonal* or alternatively the *secondary diagonal* (Nelsen, 2006, pp. 12 and 16) of copula $\mathbf{C}(u, v)$. The primary diagonal is defined as

$$\delta_{\mathbf{C}}(t) = \mathbf{C}(t, t),$$

and the secondary diagonal is defined as

$$\delta_{\mathbf{C}}^*(t) = \mathbf{C}(t, 1 - t).$$

Plotting is provided by this function because the diagonals are such important visual attributes of a copula. This function computes whole diagonals. If individual values are desired, then users are asked to use function calls along the diagonal such as `COP(0.25, 0.25, cop=P)` for the primary diagonal and `COP(0.25, 1-0.25, cop=P)` for the secondary diagonal, where for both examples the *independence copula* ($uv = \Pi$; **P**) was chosen for purposes of clarification.

The $\delta_{\mathbf{C}}(t)$ is related to order statistics of the multivariate sample (here bivariate) (Durante and Sempi, 2015, p. 68). The probability for the maxima is $\Pr[\max(u, v) \leq t] = \mathbf{C}(t, t) = \delta_{\mathbf{C}}(t)$ and the probability for the minima is $\Pr[\min(u, v) \leq t] = 2t - \delta_{\mathbf{C}}(t)$.

Usage

```

diagCOP(cop=NULL, para=NULL, secondary=FALSE,
        ploton=TRUE, lines=TRUE, delt=0.005, ...)

```

Arguments

cop	A copula function;
para	Vector of parameters, if needed, to pass to the copula;
secondary	A logical to toggle the secondary diagonal;
ploton	A logical to toggle on the plot;
lines	Draw the lines of diagonal to the current device;
delt	The increment of the diagonal curve to plot, defaults to 0.5-percent intervals, which should be small enough to resolve fine curvature for many copulas in practice; and
...	Additional arguments to pass to the plot() and lines() functions in R.

Value

An R list of the t values, $\delta_C(t, t)$ (primary) or $\delta_C^*(t, 1 - t)$ (secondary diagonal), along with a tag as to which diagonal is returned.

Author(s)

W.H. Asquith

References

Durante, F., and Sempi, C., 2015, Principles of copula theory: Boca Raton, CRC Press, 315 p.
 Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[diagCOPatf](#), [COP](#), [sectionCOP](#)

Examples

```
## Not run:
# The primary diagonal of the W, P, M, and PSP copulas on the same plot
D <- diagCOP(cop=W, lwd=2)
D <- diagCOP(cop=P, lty=2, ploton=FALSE)
D <- diagCOP(cop=M, col=2, ploton=FALSE)
D <- diagCOP(cop=PSP, col=3, ploton=FALSE)
mtext("PRIMARY DIAGONAL OF SIMPLE COPULAS") # four primary diagonals
## End(Not run)

## Not run:
# The secondary diagonal of the W, P, M, and PSP copulas on the same plot
D <- diagCOP(cop=W, lwd=2, secondary=TRUE)
D <- diagCOP(cop=P, lty=2, secondary=TRUE, ploton=FALSE)
D <- diagCOP(cop=M, col=2, secondary=TRUE, ploton=FALSE)
D <- diagCOP(cop=PSP, col=3, secondary=TRUE, ploton=FALSE)
mtext("SECONDARY DIAGONAL OF SIMPLE COPULAS") # four secondary diagonals
## End(Not run)
```

Description

Compute a numerical root along the *primary diagonal* (Nelsen, 2006, pp. 12 and 16) of copula $\mathbf{C}(u, v) = F = \mathbf{C}(t, t)$ having joint probability F . The diagonals treat the nonexceedance probabilities u and v as equals ($u = v = t$). The primary diagonal is defined for a joint nonexceedance probability t as

$$F = \mathbf{C}(t, t) \rightarrow t = \delta_{\mathbf{C}}^{(-1)}(f),$$

where the function solves for t . Examples using the concept behind diagCOPatf are available under `duCOP` and `jointCOP`, thus the diagCOPatf function can be also called by either `jointCOP` and `joint.curvesCOP`. Internally, the function uses limits of the root finder that are not equal to the anticipated interval $[0, 1]$, but equal to “small” (see description for argument interval). The function does trap for $f = 0$ by returning zero and $f = 1$ by returning unity.

Usage

```
diagCOPatf(f, cop=NULL, para=NULL, interval=NULL, silent=TRUE, verbose=FALSE,
           tol=.Machine$double.eps/10, ...)
diagCOPinv(f, cop=NULL, para=NULL, interval=NULL, silent=TRUE, verbose=FALSE,
           tol=.Machine$double.eps/10, ...)
```

Arguments

<code>f</code>	Joint probability values as a nonexceedance probability F for which to compute the root t ;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters, if needed, to pass to the copula;
<code>interval</code>	An optional interval for the root search. The default is <code>interval=c(lo, 1-lo)</code> for <code>lo=.Machine\$double.eps</code> because of difficulties for an interval on $[0, 1]$;
<code>silent</code>	The argument of the same name given over to <code>try()</code> wrapping the <code>uniroot()</code> operation;
<code>verbose</code>	If TRUE then the whole output of the numerical root is returned using only the first value provided by argument <code>f</code> ;
<code>tol</code>	The tolerance to pass to <code>uniroot</code> . The default here is much smaller than the default of the <code>uniroot()</code> function in R because of possibility that diagCOPatf would be used at extremely large nonexceedance probabilities; and
<code>...</code>	Additional arguments to pass.

Value

An R list of the root by the uniroot() function in R is returned if verbose is TRUE, otherwise the roots (diagonal inverses) for t are returned, and if an individual inverse operation fails, then a NA is returned instead.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[diagCOP](#), [jointCOP](#), [joint.curvesCOP](#)

Examples

```
diagCOPatf(0.67, cop=PSP) # 0.8023879
diagCOPatf(0.99, cop=M)  # 0.99 (now see the example below)

## Not run:
# Several functions from the lmomco package are needed.
# Suppose we have two phenomena with these log10 L-moments:
lmrA <- lmomco::vec2lmom(c(3.97, 0.485, -0.1178, 0.06857))
lmrB <- lmomco::vec2lmom(c(3.77, 0.475, -0.1377, 0.08280))
# Suppose we think that the Gumbel-Hougaard copula is appropriate with a Tau=0.45
Tau <- 0.45 # Kendall Tau between A and B.
# Suppose that the F=0.99 for either A and B provides a common risk level when they
# are considered in isolation. But what if A and B are rivers that join and joint
# FF=0.99 at their union is of interest?
FF <- 0.99
parA <- lmomco::lmom2par(lmrA, type="kap")
parB <- lmomco::lmom2par(lmrB, type="kap")
EventA <- lmomco::qlmomco(FF, parA)
EventB <- lmomco::qlmomco(FF, parB)
ApB <- 10^(EventA) + 10^(EventB) # Purely an additive conceptualization
# The FF=0.99 event is assumed to occur simultaneously on both streams, which is
# equivalent to saying that the correlation between the two is absolute 1-to-1.

# Now consider including the association as measured by Kendall Tau:
Fjoint <- diagCOPatf(FF, cop=GHCop, para=GHCop(tau=Tau)$para)
EventAj <- lmomco::qlmomco(Fjoint, parA)
EventBj <- lmomco::qlmomco(Fjoint, parB)
AcB <- 10^(EventAj) + 10^(EventBj) # Joint probability 0.99 at the union

# Now consider the association if the rivers are INDEPENDENT:
Fjoint <- diagCOPatf(FF, cop=GHCop, para=GHCop(tau=0)$para)
EventAj <- lmomco::qlmomco(Fjoint, parA)
EventBj <- lmomco::qlmomco(Fjoint, parB)
AiB <- 10^(EventAj) + 10^(EventBj) # Joint probability 0.99 at the union
```

```
# ApB = 312,000 # The perfectly simultaneous addition makes too little.
# AcB = 323,000 # The copula preserves at least the known association.
# AiB = 330,000 # The independence conceptualization makes too much.
## End(Not run)
```

duCOP

The Dual of a Copula Function

Description

Compute the *dual of a copula (function)* from a copula (Nelsen, 2006, pp. 33–34), which is defined as

$$\Pr[U \leq v \text{ or } V \leq v] = \tilde{\mathbf{C}}(u, v) = u + v - \mathbf{C}(u, v),$$

where $\tilde{\mathbf{C}}(u, v)$ is the dual of a copula and u and v are nonexceedance probabilities. The dual of a copula is the expression for the probability that either $U \leq u$ **or** $V \leq v$, which is unlike the *co-copula (function)* (see [coCOP](#)) that provides $\Pr[U > u \text{ or } V > v]$. The dual of a copula is a function and not in itself a copula. The dual of the *survival copula* ([surCOP](#)) is the *co-copula (function)* ([coCOP](#)). Some rules of copulas mean that

$$\hat{\mathbf{C}}(u', v') + \tilde{\mathbf{C}}(u, v) = 1,$$

where $\hat{\mathbf{C}}(u', v')$ is the survival copula in terms of exceedance probabilities u' and v' or in **copBasic** code that the functions [surCOP](#) + duCOP equal unity.

The function duCOP gives “protection” against simultaneous (concurrent or dual) risk by failure if and only if failure is caused (defined) by both hazard sources U and V being by themselves responsible for failure. Expressing this in terms of an annual probability of occurrence (q), one has

$$q = 1 - \Pr[U \leq v \text{ or } V \leq v] = 1 - \tilde{\mathbf{C}}(u, v) \text{ or}$$

in R code `q <- 1 - duCOP(u, v)`. So, as a mnemonic: *A dual of a copula is the probability of nonexceedance if the hazard sources must **dual** (concur, link, pair, twin, twain) between each other to cause failure.* An informative graphic is shown within [copBasic-package](#).

Usage

```
duCOP(u, v, cop=NULL, para=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula; and
<code>...</code>	Additional arguments to pass (such as parameters, if needed, for the copula in the form of a list.

Value

Value(s) for the dual of a copula are returned.

Note

There can be confusion in the interpretation and implementation of the **or** condition of *joint probability* provided by $\tilde{C}(u, v)$. Two types of **or**'s seemingly exist depending on one's concept of the meaning of "or." To start, there is the "either or both" conceptualization (**joint or**) that encompasses either "event" (say a loss) of importance for random variables U and V as well as the **joint and** conditions where both variables simultaneously are generating an event of importance.

Let us continue by performing a massive simulation for the $\text{PSP}(u, v)$ copula (**PSP**) and set an either event standard on the margins as 10 percent for an arbitrary starting point. The **PSP** has positive association with lower tail dependency, and the example here considers the left tail as the risk tail.

```
Event <- 0.1; nn <- 100000; set.seed(9238)
UV <- simCOP(n=nn, cop=PSP, graphics=FALSE) # 1E5 realizations
```

Next, let us step through counting and then make theoretical comparisons using copula theory. The **joint and** condition as nonexceedances is

```
ANDs <- length(UV$U[UV$U <= Event & UV$V <= Event]) / nn
ANDt <- COP(Event, Event, cop=PSP)
message( "Joint AND by simulation = ", round(ANDs, digits=5),
        "\n    Joint AND by theory = ", round(ANDt, digits=5))
# ANDs = 0.05348 and ANDt = 0.05263 (numerical congruence)
```

where it is obvious that the simulations and theory estimate about the same **joint and** condition. Now, the **joint or** condition as nonexceedances is

```
ORs <- length(UV$U[UV$U <= Event | UV$V <= Event]) / nn
ORt <- duCOP(Event, Event, cop=PSP)
message( "Joint OR by simulation = ", round(ORs, digits=5),
        "\n    Joint OR by theory = ", round(ORt, digits=5))
# ORs = 0.14779 and ORt = 0.14737 (numerical congruence)
```

where it is obvious that the simulations and theory estimate about the same **joint or** condition. Finally, the joint **mutually exclusive or** condition as nonexceedances is

```
eORs <- length((UV$U[(UV$U <= Event | UV$V <= Event) &
                    ! (UV$U <= Event & UV$V <= Event)])) / nn
eORt <- ORt - ANDt # theoretical computation
message( "Joint exclusive OR by simulation = ", round(eORs, digits=5),
        "\n    Joint exclusive OR by theory = ", round(eORt, digits=5))
# eORs = 0.09431 and eORt = 0.09474 (numerical congruence)
```

where it is obvious that the simulations and theory estimate about the same joint **mutually exclusive or** condition, and where it is shown that the prior two theoretical joint probabilities can be subtracted from each to yield the **mutually exclusive or** condition.

Let us then play out a scenario in which it is judged that of the events causing damage that the simultaneous occurrence is worse but that engineering against about 5 percent of events not occurring at the same time represents the most funding available. Using numerical methods, it is possible to combine $\tilde{C}$ and C and assume equal marginal risk in U and V as the following list shows:

```
"designf" <- function(t) { # a one-off function just for this example
  duCOP(t, t, cop=PSP) - COP(t, t, cop=PSP) - 5/100 # 5 percent
}
dThres <- uniroot(designf, c(.Machine$double.eps,0.5))$root
```

where the uniroot function performs the optimization and the .Machine\$double.eps value is used because the **PSP** is NaN for zero probability. (It is unity for unity marginal probabilities.)

The design threshold on the margins then is $dThres \approx 0.05135$. In other words, the designThres is the marginal probability that results in about 5 percent of events not occurring at the same time. Then considering the simulated sample and counting the nonexceedances by code one achieves:

```
Damage      <- length( UV$U[ UV$U <= dThres | UV$V <= dThres ])
SimDamage    <- length( UV$U[ UV$U <= dThres & UV$V <= dThres ])
NonSimDamage <- length((UV$U[(UV$U <= dThres | UV$V <= dThres) &
                           ! (UV$U <= dThres & UV$V <= dThres)]))
message( "          Damaging Events (sim.) = ", Damage,
  "\n    Simultaneous damaging events (sim.) = ", SimDamage,
  "\n Nonsimultaneous damaging events (sim.) = ", NonSimDamage)
```

but also the theoretical expectations are readily computed using copula theory:

```
tDamage      <- as.integer(duCOP(dThres, dThres, cop=PSP) * nn)
tSimDamage    <- as.integer( COP(dThres, dThres, cop=PSP) * nn)
tNonSimDamage <- tDamage - tSimDamage
message( "          Damaging Events (theory) = ", tDamage,
  "\n    Simultaneous damaging events (theory) = ", tSimDamage,
  "\n Nonsimultaneous damaging events (theory) = ", tNonSimDamage)
```

The counts from the former listing are 7,670; 2,669; and 5,001, whereas the respective counts from the later listing are 7,635; 2,635; and 5,000. Numerical congruency in the counts thus exists.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[COP](#), [coCOP](#), [surCOP](#), [jointCOP](#), [joint.curvesCOP](#)

Examples

```

u <- runif(1); t <- runif(1)
duCOP(cop=W,u,t)    # joint or probability for perfect negative dependence
duCOP(cop=P,u,t)    # joint or probability for perfect independence
duCOP(cop=M,u,t)    # joint or probability for perfect positive dependence
duCOP(cop=PSP,u,t)  # joint or probability for some positive dependence

# Next demonstrate COP + duCOP = unity.
"M0cop.formula" <- function(u,v, para=para, ...) {
  alpha <- para[1]; beta <- para[2]; return(min(v*u^(1-alpha), u*v^(1-beta)))
}
"M0cop" <- function(u,v, ...) { asCOP(u,v, f=M0cop.formula, ...) }

u <- 0.2; v <- 0.75; ab <- c(1.5, 0.3)
surCOP(1-u,1-v, cop=M0cop, para=ab) + duCOP(u,v, cop=M0cop, para=ab) # UNITY

# See extended code listings and discussion in the Note section

```

EMPIRcop

The Bivariate Empirical Copula

Description

The *bivariate empirical copula* (Nelsen, 2006, p. 219) for a bivariate sample of length n is defined for random variables X and Y as

$$\mathbf{C}_n\left(\frac{i}{n}, \frac{j}{n}\right) = \frac{\text{number of pairs } (x, y) \text{ with } x \leq x_{(i)} \text{ and } y \leq y_{(j)}}{n},$$

where $x_{(i)}$ and $y_{(i)}$, $1 \leq i, j \leq n$ or expressed as

$$\mathbf{C}_n\left(\frac{i}{n}, \frac{j}{n}\right) = \frac{1}{n} \sum_{i=1}^n \mathbf{1}\left(\frac{R_i}{n} \leq u_i, \frac{S_i}{n} \leq v_i\right),$$

where R_i and S_i are ranks of the data for U and V , and $\mathbf{1}(\cdot)$ is an *indicator function* that score 1 if condition is true otherwise scoring zero. Using more generic notation, the empirical copula can be defined by

$$\mathbf{C}_n(u, v) = \frac{1}{n} \sum_{i=1}^n \mathbf{1}(u_i^{\text{obs}} \leq u_i, v_i^{\text{obs}} \leq v_i),$$

where u^{obs} and v^{obs} are thus some type of nonparametric nonexceedance probabilities based on counts of the underlying data expressed in probabilities.

Hazen Empirical Copula—The “Hazen form” of the empirical copula is

$$\mathbf{C}_n^{\mathcal{H}}(u, v) = \frac{1}{n} \sum_{i=1}^n \mathbf{1}\left(\frac{R_i - 0.5}{n} \leq u_i, \frac{S_i - 0.5}{n} \leq v_i\right),$$

which can be triggered by `ctype="hazen"`. This form is named for this package because of direct similarity of the *Hazen plotting position* to the above definition. Joe (2014, pp. 247–248) uses the Hazen form. Joe continues by saying “[the] adjustment of the uniform score $[(R - 0.5)/n]$ could be done in an alternative form, but there is [asymptotic] equivalence[, and that] C_n^H puts mass of n^{-1} at the tuples $([r_{i1} - 0.5]/n, \dots, [r_{id} - 0.5]/n)$ for $i = 1, \dots, n$.” A footnote by Joe (2014) says that “the conversion $[R/(n + 1)]$ is commonly used for the empirical copula.” This later form is the “Weibull form” described next. Joe’s preference for the Hazen form is so that the sum of squared normal scores is closer to unity for large n than such a sum would be attained using the Weibull form.

Weibull Empirical Copula—The “Weibull form” of the empirical copula is

$$C_n^W(u, v) = \frac{1}{n} \sum_{i=1}^n \mathbf{1}\left(\frac{R_i}{n+1} \leq u_i, \frac{S_i}{n+1} \leq v_i\right),$$

which can be triggered by `ctype="weibull"`. This form is named for this package because of direct similarity of the *Weibull plotting position* to the definition, and this form is the default (see argument description).

Bernstein Empirical Copula—The empirical copula can be extended nonparametrically as the *Bernstein empirical copula* (Hernández-Maldonado *et al.*, 2012) and is formulated as

$$C_n^B(u, v; \eta) = \sum_{i=1}^n \sum_{j=1}^n C_n\left(\frac{i}{n}, \frac{j}{n}\right) \times \eta(i, j; u, v),$$

where the individual *Bernstein weights* $\eta(i, j)$ for the k th paired value of the u and v vectors are

$$\eta(i, j; u, v) = \binom{n}{i} u^i (1-u)^{n-i} \times \binom{n}{j} u^j (1-u)^{n-j}.$$

The Bernstein extension, albeit conceptually pure in its shuffling by binomial coefficients and left- and right-tail weightings, is quite CPU intensive as inspection of the equations above indicates a nest of four `for()` loops in R. (The native R code of **copBasic** uses the `sapply()` function in R liberally for substantial but not a blazing fast speed increase.) The Bernstein extension results in a smoother surface of the empirical copula and can be triggered by `ctype="bernstein"`. Remark, Hernández-Maldonado *et al.* (2024) provide a fast algorithm for the Bernstein smoothing—this implemented in [EMPIRgrid_fast](#).

Checkerboard Empirical Copula—A simple smoothing to the empirical copula is the *checkerboard empirical copula* (Segers *et al.*, 2017) that has been adapted from the **copula** package. It is numerically intensive like the Bernstein and possibly of limited usefulness for large sample sizes. The checkerboard extension can be triggered by `ctype="checkerboard"` and is formulated as

$$C_n^\#(U) = \frac{1}{n+o} \sum_{i=1}^n \prod_{j=1}^d \min[\max[nU_j - R_{i,j}^{(n)} + 1, 0], 1],$$

where U is a $d = 2$ column matrix of u and v , R is a rank function, and o is an offset term on $[0, 1]$.

The *empirical copula frequency* can be defined (Nelson, 2006, p. 219) as

$$c_n(u, v) = C_n\left(\frac{i}{n}, \frac{j}{n}\right) - C_n\left(\frac{i-1}{n}, \frac{j}{n}\right) - C_n\left(\frac{i}{n}, \frac{j-1}{n}\right) + C_n\left(\frac{i-1}{n}, \frac{j-1}{n}\right).$$

Remembering that the several asymptotically equivalent definitions of the empirical copula do converge as n becomes large, we can show small-sample congruence between the empirical copula from the **copula** package and **copBasic** to the Weibull version of EMPIRcop because of **copula::pobs()** is itself Weibull:

```
uv <- as.matrix( simCOP(50, cop=P) )
ctype <- "weibull"; # Try the other ctypes supported by EMPIRcop
a <- EMPIRcop(uv[,1], uv[,2], para=as.data.frame(uv), ctype=ctype)
b <- copula::C.n(uv, uv, offset=0) # if ctype="weibull" equal values
plot(a, b, xlab=paste0('copBasic EMPIRcop(ctype=" ', ctype, ' ")'),
      ylab="copula::C.n"); abline(0,1, col="red")
```

Usage

```
EMPIRcop(u, v, para=NULL,
         ctype=c("weibull", "hazen", "1/n", "bernstein", "checkerboard"),
         bernprogress=FALSE, checkerboard.offset=0, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the U-statistics of the data (see Examples). Alternatively, <code>para</code> can be a list holding a <code>para</code> as would be done if it were a vector, but arguments <code>bernstein</code> and <code>bernprogress</code> can be optionally included—this feature is provided so that the Bernstein refinement can be controlled within the context of other functions calling EMPIRcop such as by level.curvesCOP ;
<code>ctype</code>	An alternative means for triggering the definition of C_n , C_n^W (default), C_n^H , C_n^B , or $C_n^\#$. This argument of the same name is also used by blomCOP ;
<code>bernprogress</code>	The Bernstein copula extension is CPU intensive(!), so a splash counter is pushed to the console via the <code>message()</code> function in R so as to not discourage the user;
<code>checkerboard.offset</code>	A scaling of the ratio <code>sum(...)/(n+offset)</code> for the checkerboard empirical copula; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Note

Not all theoretical measures of copula dependence (both measures of association and measures of asymmetry), which use numerical integration by the `integrate()` function in R, can be used for all empirical copulas because of “divergent” integral errors; however, examples using *Hoeffding Phi* (Φ_C ; [hoefCOP](#)) and shown under **Examples**. Other measures of copula dependence include

`blomCOP`, `footCOP`, `giniCOP`, `rhoCOP`, `tauCOP`, `wolfCOP`, `joeskewCOP`, and `uvlmoms`. Each of these measures fortunately has a built-in sample estimator.

It is important to distinguish between a sample estimator and the estimation of the measure using the empirical copula itself via the `EMPIRCOP` function. The sample estimators (triggered by the `as.sample` arguments for the measures) are reasonably fast and numerically preferred over using the empirical copula. Further, the generally slow numerical integrations for the theoretical definitions of these copula measures might have difficulties. Limited testing, however, suggests prevalence of numerical integration not erroring using the Bernstein extension of the empirical copula, which must be a by-product of the enhanced and sufficient smoothness for the R default numerical integration to succeed. Many of the measures have brute option for a brute-force numerical integration on a regular grid across the empirical copula—these are slow but should not trigger errors. As a general rule, users should still use the sample estimators instead.

Author(s)

W.H. Asquith

References

- Hernández-Maldonado, V.M., Díaz-Viera, M., and Erdely, A., 2012, A joint stochastic simulation method using the Bernstein copula as a flexible tool for modeling nonlinear dependence structures between petrophysical properties: *Journal of Petroleum Science and Engineering*, v. 90–91, pp. 112–123, [doi:10.1016/j.petrol.2012.04.018](https://doi.org/10.1016/j.petrol.2012.04.018).
- Hernández-Maldonado, V.M., Erdely, A., Díaz-Viera, M., and Rios, L., 2024, Fast procedure to compute empirical and Bernstein copulas: *Applied Mathematics and Computation*, v. 477, article 128827, 14 p., [doi:10.1016/j.amc.2024.128827](https://doi.org/10.1016/j.amc.2024.128827).
- Nelsen, R.B., 2006, *An introduction to copulas*: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, *Extremes in Nature—An approach using copulas*: Springer, 289 p.
- Segers, J., Sibuya, M., and Tsukahara, H., 2017, The empirical beta copula: *Journal of Multivariate Analysis*, v. 155, pp. 35–51, [doi:10.1016/j.jmva.2016.11.010](https://doi.org/10.1016/j.jmva.2016.11.010).

See Also

`diagCOP`, `level.curvesCOP`, `simCOP`

Examples

```
## Not run:
set.seed(62)
EMPIRCOP(0.321,0.78, para=simCOP(n=90, cop=N4212cop,
                                para=2.32, graphics=FALSE)) # [1] 0.3222222
N4212cop(0.321,0.78, para=2.32)                               # [1] 0.3201281
## End(Not run)

## Not run:
set.seed(62) # See note below about another seed to try.
psp <- simCOP(n=34, cop=PSP, ploton=FALSE, points=FALSE) * 150
# Pretend psp is real data, the * 150 is to clearly get into an arbitrary unit system.
```

```

# The sort=FALSE is critical in the following two calls. Although the Weibull
# plotting positions are chosen, internally EMPIRcop uses ranks, but the model
# here is to imagine one having a sample in native units of the random variables
# and then casting them into probabilities for other purposes.
fakeU <- lmomco::pp(psp[,1], sort=FALSE) # Weibull plotting position i/(n+1)
fakeV <- lmomco::pp(psp[,2], sort=FALSE) # Weibull plotting position i/(n+1)
uv <- data.frame(U=fakeU, V=fakeV); # our U-statistics

# The next four values should be very close if n above were say 1000, but the
# ctype="bernstein" should not be used if n >> 34 because of inherently long runtime.
PSP(0.4,0.6)           # 0.3157895 (compare to listed values below)

# Two seeds are shown so that the user can see that depending on the distribution
# of the values given by para that different coincidences of which method is
# equivalent to another exist.
# For set.seed(62) --- "hazen" == "weibull" by coincidence
#   "hazen"      --> 0.3529412
#   "weibull"    --> 0.3529412
#   "1/n"        --> 0.3235294
#   "bernstein"  --> 0.3228916
# For set.seed(85) --- "1/n" == "hazen" by coincidence
#   "hazen"      --> 0.3529412
#   "weibull"    --> 0.3823529
#   "1/n"        --> 0.3529412
#   "bernstein"  --> 0.3440387

# For set.seed(62) --- not all measures of association can be used for all
# empirical copulas because of 'divergent' integral errors, but this is an example
# for Hoeffding Phi. These computations are CPU intensive, esp. Bernstein.
hoefCOP(as.sample=TRUE, para=uv) # (sample estimator is fast) # 0.4987755
hoefCOP(cop=EMPIRcop, para=uv, ctype="hazen") # 0.5035348
hoefCOP(cop=EMPIRcop, para=uv, ctype="weibull") # 0.4977145
hoefCOP(cop=EMPIRcop, para=uv, ctype="1/n") # 0.4003646
hoefCOP(cop=EMPIRcop, para=uv, ctype="bernstein") # 0.4563724
hoefCOP(cop=EMPIRcop, para=uv, ctype="checkerboard") # 0.4952427
## End(Not run)

# All other example suites shown below are dependent on the pseudo-data in the
# variable uv. It is suggested to not run with a sample size much larger than the
# above n=34 if the Bernstein comparison is intended (wanted) simply because of
# lengthy(!) run times, but the n=34 does provide a solid demonstration how the
# level curves for Bernstein weights are quite smooth.

## Not run:
# Now, let us construct as many as three sets of level curves to the sample
# resided in the uv sample from above using the PSP copula.
level.curvesCOP(cop=PSP); # TRUE, parametric, fast, BLACK CURVES

# Empirical copulas can consume lots of CPU.
# RED CURVES, if n is too small, uniroot() errors might be triggered and likely
# will be using the sample size of 34 shown above.
level.curvesCOP(cop=EMPIRcop, para=uv, delu=0.03, col=2, ploton=FALSE)

```

```

# GREEN CURVES (large CPU commitment)
# Bernstein progress is uninformative because level.curvesCOP() has taken over control.
bpara <- list(para=uv, ctype="bernstein", bernprogress=FALSE)
level.curvesCOP(cop=EMPIRcop, para=bpara, delu=0.03, col=3, ploton=FALSE)
# The delu is increased for faster execution but more important,
# notice the greater smoothness of the Bernstein refinement.
## End(Not run)

## Not run:
# Experimental from R Graphics by Murrell (2005, p.112)
"trans3d" <-
function(x,y,z, pmat) {
  tmat <- cbind(x,y,z,1) %*% pmat
  return(tmat[,1:2] / tmat[,4])
}

the.grid <- EMPIRgrid(para=uv, ctype="checkerboard")
the.diag <- diagCOP(cop=EMPIRcop, para=uv, ploton=FALSE, lines=FALSE)

the.persp <- persp(the.grid$empcop, theta=-25, phi=20,
  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")
the.trace <- trans3d(the.diag$t, the.diag$t, the.diag$diagcop, the.persp)
lines(the.trace, lwd=2, col=2) # The diagonal of the copula

# The following could have been used as an alternative to call persp()
the.persp <- persp(x=the.grid$u, y=the.grid$v, z=the.grid$empcop, theta=-25, phi=20,
  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")
lines(the.trace, lwd=2, col=2) # The diagonal of the copula #
## End(Not run)

```

EMPIRcopdf

*Data Frame Representation of the Bivariate Empirical Copula***Description**

Generate an R data.frame representation of the *bivariate empirical copula* (Salvadori *et al.*, 2007, p. 140) using the coordinates as preserved in the raw data in the parameter object of the bivariate empirical copula.

Usage

```
EMPIRcopdf(para=NULL, ...)
```

Arguments

para	A vector (single element) of parameters—the U-statistics of the data (see example) to pass to EMPIRcop ; and
...	Additional arguments to pass to EMPIRcop .

Value

An R data.frame of u , v , and $C_n(u, v)$ values of the bivariate empirical copula is returned.

Author(s)

W.H. Asquith

References

Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, *Extremes in Nature—An approach using copulas*: Springer, 289 p.

See Also

[EMPIRcop](#)

Examples

```
## Not run:
psp <- simCOP(n=39, cop=PSP, ploton=FALSE, points=FALSE) * 150
# Pretend psp is real data, the * 150 is to clearly get into an arbitrary unit system.

# The sort=FALSE is critical in the following two calls to pp() from lmomco.
fakeU <- lmomco::pp(psp[,1], sort=FALSE) # Weibull plotting position i/(n+1)
fakeV <- lmomco::pp(psp[,2], sort=FALSE) # Weibull plotting position i/(n+1)
uv <- data.frame(U=fakeU, V=fakeV) # our U-statistics

empcop <- EMPIRcopdf(para=uv)
plot(empcop$u, empcop$v, cex=1.75*empcop$empcop, pch=16,
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
# Dot size increases with joint probability (height of the copulatic surface).
points(empcop$u, empcop$v, col=2) # red circles
## End(Not run)
```

Description

Generate a gridded representation of the *bivariate empirical copula* (see [EMPIRcop](#), Salvadori *et al.*, 2007, p. 140). Herein, the $\check{C}_n$ (double-dots) are used to denote gridding of the copula. This function has the primary intention of supporting 3-D renderings or 2-D images of the *copulatic surface*, but many empirical copula functions in **copBasic** rely on the grid of the empirical copula—unlike the functions that support parametric copulas. The algorithm within EMPIRgrid is slow because of nested loops, but a vastly faster version with some tweaks (not having an explicit `de1uv`) is available under [EMPIRgrid_fast](#).

Usage

```
EMPIRgrid(para=NULL, deluv=0.05, gridonly=FALSE, verbose=FALSE, ...)
```

Arguments

para	A matrix of u and v vectors as an alternative to uv for syntax compatibility for other situations in copBasic ;
deluv	A delta value of the both the u and v axes (grid edges) for empirical copula estimation by the EMPIRcop function;
gridonly	A logical to return the empcop component ($\ddot{C}_n(u, v)$) passed through ... the ctype for EMPIRcop ;
verbose	A logical controlling whether the progress during grid building is to be shown; and
...	Additional arguments to pass to EMPIRcop .

Value

A matrix is returned ($\ddot{C}_n(u, v)$), if `gridonly` is true. Otherwise a list of the u , v , and $\ddot{C}_n(u, v)$ or other definitions (ctype for **EMPIRcop**) is returned. The list will also contain the `deluv` used to generate the grid and if the user passed through ... the ctype for **EMPIRcop**, then that is preserved, otherwise set at "unknown". The list is returned as classes of "empirical.copula.grid" and "list". This classification might be used in **copBasic** to check for a grid in other functions.

Note

The extensive suite of examples is included here because the various ways that algorithms involving empirical copulas can be tested. The figures also provide excellent tools for education on copulas.

Author(s)

W.H. Asquith

References

Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, *Extremes in Nature—An approach using copulas*: Springer, 289 p.

See Also

[EMPIRcop](#), [EMPIRgrid_fast](#), [EMPIRgrid_lkup](#), [EMPIRcopdf](#)

Examples

```
## Not run:
# EXAMPLE 1:
psp <- simCOP(n=490, cop=PSP, ploton=FALSE, points=FALSE) * 150
# Pretend psp is real data, the * 150 is to clearly get into an arbitrary unit system.

# The sort=FALSE is critical in the following two calls to pp() from lmomco.
```

```

fakeU <- lmomco::pp(ppsp[,1], sort=FALSE) # Weibull plotting position i/(n+1)
fakeV <- lmomco::pp(ppsp[,2], sort=FALSE) # Weibull plotting position i/(n+1)
uv <- data.frame(U=fakeU, V=fakeV) # our U-statistics

# The follow function is used to make 3-D --> 2-D transformation
# From R Graphics by Murrell (2005, p.112)
"trans3d" <- # back slashes seem needed for the package
function(x,y,z, pmat) { # for user manual building but bad syntax
  tmat <- cbind(x,y,z,1) %*% pmat # because remember the percent sign is a
  return(tmat[,1:2] / tmat[,4]) # a comment character in LaTeX.
}

the.grid <- EMPIRgrid(para=uv)
cop.diag <- diagCOP(cop=EMPIRcop, para=uv, ploton=FALSE, lines=FALSE)
empcop <- EMPIRcopdf(para=uv) # data.frame of all points

# EXAMPLE 1: PLOT NUMBER 1
the.persp <- persp(the.grid$empcop, theta=-25, phi=20,
  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")

# EXAMPLE 1: PLOT NUMBER 2 (see change in interaction with variable 'the.grid')
the.persp <- persp(x=the.grid$u, y=the.grid$v, z=the.grid$empcop, theta=-25, phi=20,
  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")

the.diag <- trans3d(cop.diag$t, cop.diag$t, cop.diag$diagcop, the.persp)
lines(the.diag, lwd=4, col=3, lty=2)

points(trans3d(empcop$u, empcop$v, empcop$empcop, the.persp),
  col=rgb(0,1-sqrt(empcop$empcop),1,sqrt(empcop$empcop)), pch=16)
# the sqrt() is needed to darken or enhance the colors

S <- sectionCOP(cop=PSP, 0.2, ploton=FALSE, lines=FALSE)
thelines <- trans3d(rep(0.2, length(S$t)), S$t, S$seccop, the.persp)
lines(thelines, lwd=2, col=6)
S <- sectionCOP(cop=PSP, 0.2, ploton=FALSE, lines=FALSE, dercop=TRUE)
thelines <- trans3d(rep(0.2, length(S$t)), S$t, S$seccop, the.persp)
lines(thelines, lwd=2, col=6, lty=2)

S <- sectionCOP(cop=PSP, 0.85, ploton=FALSE, lines=FALSE, wrtV=TRUE)
thelines <- trans3d(S$t, rep(0.85, length(S$t)), S$seccop, the.persp)
lines(thelines, lwd=2, col=2)
S <- sectionCOP(cop=PSP, 0.85, ploton=FALSE, lines=FALSE, dercop=TRUE)
thelines <- trans3d(S$t, rep(0.85, length(S$t)), S$seccop, the.persp)
lines(thelines, lwd=2, col=2, lty=2)

empder <- EMPIRgridder(empgrid=the.grid)
thelines <- trans3d(rep(0.2, length(the.grid$v)), the.grid$v, empder[,3], the.persp)
lines(thelines, lwd=4, col=6) #
## End(Not run)

## Not run:
# EXAMPLE 2:
# An asymmetric example to demonstrate that the grid is populated with the

```

```

# correct orientation---meaning U is the horizontal and V is the vertical.

# EXAMPLE 2: PLOT NUMBER 1 # See the asymmetry
uv <- simCOP(1000, cop=MOCop, para=c(0.4,0.9)) # The parameters cause asymmetry.
mtext("Simulation from a defined Marshall-Olkin Copula")
the.grid <- EMPIRgrid(para=uv, deluv=0.025)

# EXAMPLE 2: PLOT NUMBER 2
# The second plot by image() will show a "hook" of sorts along the singularity.
image(the.grid$empcop, col=terrain.colors(40)) # Second plot is made
mtext("Image of gridded Empirical Copula")

# EXAMPLE 2: PLOT NUMBER 3
empcop <- EMPIRcopdf(para=uv) # data.frame of all points
# The third plot is the 3-D version overlain with the data points.
the.persp <- persp(x=the.grid$u, y=the.grid$v, z=the.grid$empcop, theta=240, phi=40,
                  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")
points(trans3d(empcop$u, empcop$v, empcop$empcop, the.persp),
       col=rgb(0,1-sqrt(empcop$empcop),1,sqrt(empcop$empcop)), pch=16)
mtext("3-D representation of gridded empirical copula with data points")

# EXAMPLE 2: PLOT NUMBER 4
# The fourth plot shows a simulation and the quasi-emergence of the singularity
# that of course the empirical perspective "knows" nothing about. Do not use
# the Kumaraswamy smoothing because in this case the singularity because
# too smoothed out relative to the raw empirical, but of course the sample size
# is large enough to see such things. (Try kumaraswamy=TRUE)
empsim <- EMPIRsim(n=1000, empgrid = the.grid, kumaraswamy=FALSE)
mtext("Simulations from the Empirical Copula") #
## End(Not run)

## Not run:
# EXAMPLE 3:
psp <- simCOP(n=4900, cop=PSP, ploton=FALSE, points=FALSE) * 150
# Pretend psp is real data, the * 150 is to clearly get into an arbitrary unit system.

# The sort=FALSE is critical in the following two calls to pp() from lmomco.
fakeU <- lmomco::pp(psp[,1], sort=FALSE) # Weibull plotting position i/(n+1)
fakeV <- lmomco::pp(psp[,2], sort=FALSE) # Weibull plotting position i/(n+1)
uv <- data.frame(U=fakeU, V=fakeV) # our U-statistics

# EXAMPLE 3: # PLOT NUMBER 1
deluv <- 0.0125 # going to cause long run time with large n
# The small deluv is used to explore solution quality at U=0 and 1.
the.grid <- EMPIRgrid(para=uv, deluv=deluv)
the.persp <- persp(x=the.grid$u, y=the.grid$v, z=the.grid$empcop, theta=-25, phi=20,
                  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")

S <- sectionCOP(cop=PSP, 1, ploton=FALSE, lines=FALSE)
thelines <- trans3d(rep(1, length(S$t)), S$t, S$seccop, the.persp)
lines(thelines, lwd=2, col=2)

S <- sectionCOP(cop=PSP, 0, ploton=FALSE, lines=FALSE)

```

```

thelines <- trans3d(rep(0, length(S$t)), S$t, S$seccop, the.persp)
lines(thelines, lwd=2, col=2)

S <- sectionCOP(cop=PSP, 1, ploton=FALSE, lines=FALSE, dercop=TRUE)
thelines <- trans3d(rep(1, length(S$t)), S$t, S$seccop, the.persp)
lines(thelines, lwd=2, col=2, lty=2)

S <- sectionCOP(cop=PSP, 2*deluv, ploton=FALSE, lines=FALSE, dercop=TRUE)
thelines <- trans3d(rep(2*deluv, length(S$t)), S$t, S$seccop, the.persp)
lines(thelines, lwd=2, col=2, lty=2)

empder <- EMPIRgridder(empgrid=the.grid)
thelines <- trans3d(rep(2*deluv, length(the.grid$v)), the.grid$v, empder[3,], the.persp)
lines(thelines, lwd=4, col=5, lty=2)

pdf("conditional_distributions.pdf")
ix <- 1:length(attributes(empder)$rownames)
for(i in ix) {
  u <- as.numeric(attributes(empder)$rownames[i])
  S <- sectionCOP(cop=PSP, u, ploton=FALSE, lines=FALSE, dercop=TRUE)
  # The red line is the true.
  plot(S$t, S$seccop, lwd=2, col=2, lty=2, type="l", xlim=c(0,1), ylim=c(0,1),
        xlab="V, NONEXCEEDANCE PROBABILITY", ylab="V, VALUE")
  lines(the.grid$v, empder[i,], lwd=4, col=5, lty=2) # empirical
  mtext(paste("Conditioned on U=", u, " nonexceedance probability"))
}
dev.off() #
## End(Not run)

```

EMPIRgridder

Derivatives of the Grid of the Bivariate Empirical Copula for V with respect to U

Description

Generate derivatives of V with respect to U of a gridded representation of the *bivariate empirical copula* (see [EMPIRCop](#)). This function is the empirical analog to [derCOP](#).

Usage

```
EMPIRgridder(empgrid=NULL, ...)
```

Arguments

<code>empgrid</code>	The grid from EMPIRgrid ; and
<code>...</code>	Additional arguments to pass.

Value

The gridded values of the derivatives of the bivariate empirical copula.

Author(s)

W.H. Asquith

See Also[EMPIRcop](#), [EMPIRcopdf](#), [EMPIRgrid](#), [EMPIRgridder2](#)**Examples**

```
## Not run:
para <- list(alpha=0.15, beta=0.65, cop1=PLACKETTcop, cop2=PLACKETTcop,
             para1=0.005, para2=1000)
uv <- simCOP(n=1000, cop=composite2COP, para=para)
fakeU <- lmomco::pp(uv[,1], sort=FALSE)
fakeV <- lmomco::pp(uv[,2], sort=FALSE)
uv <- data.frame(U=fakeU, V=fakeV)

"trans3d" <-                                     # back slashes seem needed for the package
function(x,y,z, pmat) {                         # for user manual building but bad syntax
  tmat <- cbind(x,y,z,1) %*% pmat               # because remember the percent sign is a
  return(tmat[,1:2] / tmat[,4])                 # a comment character in LaTeX.
}

the.grid <- EMPIRgrid(para=uv, deluv=0.1)
the.diag <- diagCOP(cop=EMPIRcop, para=uv, ploton=FALSE, lines=FALSE)
empcop <- EMPIRcopdf(para=uv) # data.frame of all points

the.persp <- persp(the.grid$empcop, theta=-25, phi=20,
                  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")
points(trans3d(empcop$u, empcop$v, empcop$empcop, the.persp),
       col=rgb(0,1-sqrt(empcop$empcop),1,sqrt(empcop$empcop)), pch=16, cex=0.75)

# Now extract the copula sections
some.lines <- trans3d(rep(0.2, length(the.grid$v)),
                     the.grid$v, the.grid$empcop[3,], the.persp)
lines(some.lines, lwd=2, col=2)
some.lines <- trans3d(the.grid$u, rep(0.6, length(the.grid$u)),
                     the.grid$empcop[7,], the.persp)
lines(some.lines, lwd=2, col=3)
some.lines <- trans3d(rep(0.7, length(the.grid$v)), the.grid$v,
                     the.grid$empcop[8,], the.persp)
lines(some.lines, lwd=2, col=6)

# Now compute some derivatives or conditional cumulative
# distribution functions
empder <- EMPIRgridder(empgrid=the.grid)
some.lines <- trans3d(rep(0.2, length(the.grid$v)), the.grid$v, empder[3,], the.persp)
lines(some.lines, lwd=4, col=2)

empder <- EMPIRgridder2(empgrid=the.grid)
some.lines <- trans3d(the.grid$u, rep(0.6, length(the.grid$u)), empder[,7], the.persp)
lines(some.lines, lwd=4, col=3)
```

```
empder <- EMPIRgridder(empgrid=the.grid)
some.lines <- trans3d(rep(0.7, length(the.grid$v)), the.grid$v, empder[8,], the.persp)
lines(some.lines, lwd=4, col=6)

# Demonstrate conditional quantile function extraction for
# the 70th percentile of U and see how it plots on top of
# the thick purple line
empinv <- EMPIRgridderinv(empgrid=the.grid)
some.lines <- trans3d(rep(0.7, length(the.grid$v)), empinv[8,],
  attributes(empinv)$colnames, the.persp)
lines(some.lines, lwd=4, col=5, lty=2)#
## End(Not run)
```

EMPIRgridder2	<i>Derivatives of the Grid of the Bivariate Empirical Copula for U with respect to V</i>
---------------	--

Description

Generate derivatives of U with respect to V of a gridded representation of the *bivariate empirical copula* (see [EMPIRcop](#)). This function is the empirical analog to [derCOP2](#).

Usage

```
EMPIRgridder2(empgrid=NULL, ...)
```

Arguments

<code>empgrid</code>	The grid from EMPIRgrid ; and
<code>...</code>	Additional arguments to pass.

Value

The gridded values of the derivatives of the bivariate empirical copula.

Author(s)

W.H. Asquith

See Also

[EMPIRcop](#), [EMPIRcopdf](#), [EMPIRgrid](#), [EMPIRgridder](#)

Examples

```
# See examples under EMPIRgridder
```

EMPIRgridderinv	<i>Derivative Inverses of the Grid of the Bivariate Empirical Copula for V with respect to U</i>
-----------------	--

Description

Generate a gridded representation of the inverse of the derivatives of the *bivariate empirical copula* of V with respect to U . This function is the empirical analog to [derCOPinv](#).

Usage

```
EMPIRgridderinv(empgrid=NULL, kumaraswamy=FALSE, dergrid=NULL, ...)
```

Arguments

empgrid	The grid from EMPIRgrid ;
kumaraswamy	A logical to trigger Kumaraswamy smoothing of the conditional quantile function;
dergrid	The results of EMPIRgridder and if left NULL then that function is called internally. There is some fragility at times in the quality of the numerical derivative and the author has provided this argument so that the derivative can be computed externally and then fed to this inversion function; and
...	Additional arguments to pass.

Value

The gridded values of the inverse of the derivative of V with respect U .

Author(s)

W.H. Asquith

See Also

[EMPIRcop](#), [EMPIRcopdf](#), [EMPIRgrid](#), [EMPIRgridder2](#)

Examples

```
## Not run:
uv <- simCOP(n=10000, cop=PSP, ploton=FALSE, points=FALSE)
fakeU <- lmomco::pp(uv[,1], sort=FALSE)
fakeV <- lmomco::pp(uv[,2], sort=FALSE)
uv <- data.frame(U=fakeU, V=fakeV)

uv.grid <- EMPIRgrid(para=uv, deluv=.1) # CPU hungry
uv.inv1 <- EMPIRgridderinv(empgrid=uv.grid)
uv.inv2 <- EMPIRgridderinv2(empgrid=uv.grid)
plot(uv, pch=16, col=rgb(0,0,0,.1), xlim=c(0,1), ylim=c(0,1),
```

```

      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
lines(qua.regressCOP(f=0.5, cop=PSP), col=2)
lines(qua.regressCOP(f=0.2, cop=PSP), col=2)
lines(qua.regressCOP(f=0.7, cop=PSP), col=2)
lines(qua.regressCOP(f=0.1, cop=PSP), col=2)
lines(qua.regressCOP(f=0.9, cop=PSP), col=2)

med.wrtu <- EMPIRqua.regress(f=0.5, empinv=uv.in1)
lines(med.wrtu, col=2, lwd=4)
qua.wrtu <- EMPIRqua.regress(f=0.2, empinv=uv.in1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.7, empinv=uv.in1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.1, empinv=uv.in1)
lines(qua.wrtu, col=2, lwd=2, lty=4)
qua.wrtu <- EMPIRqua.regress(f=0.9, empinv=uv.in1)
lines(qua.wrtu, col=2, lwd=2, lty=4)

lines(qua.regressCOP2(f=0.5, cop=PSP), col=4)
lines(qua.regressCOP2(f=0.2, cop=PSP), col=4)
lines(qua.regressCOP2(f=0.7, cop=PSP), col=4)
lines(qua.regressCOP2(f=0.1, cop=PSP), col=4)
lines(qua.regressCOP2(f=0.9, cop=PSP), col=4)

med.wrtv <- EMPIRqua.regress2(f=0.5, empinv=uv.in2)
lines(med.wrtv, col=4, lwd=4)
qua.wrtv <- EMPIRqua.regress2(f=0.2, empinv=uv.in2)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.7, empinv=uv.in2)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.1, empinv=uv.in2)
lines(qua.wrtv, col=4, lwd=2, lty=4)
qua.wrtv <- EMPIRqua.regress2(f=0.9, empinv=uv.in2)
lines(qua.wrtv, col=4, lwd=2, lty=4)#
## End(Not run)

## Not run:
# Now try a much more complex shape
para <- list(alpha=0.15, beta=0.65, cop1=PLACKETTcop, cop2=PLACKETTcop,
             para1=0.005, para2=1000)
uv <- simCOP(n=30000, cop=composite2COP, para=para)
fakeU <- lmomco::pp(uv[,1], sort=FALSE)
fakeV <- lmomco::pp(uv[,2], sort=FALSE)
uv <- data.frame(U=fakeU, V=fakeV)

uv.grid <- EMPIRgrid(para=uv, deluv=0.05) # CPU hungry
uv.in1 <- EMPIRgridderinv(empgrid=uv.grid)
uv.in2 <- EMPIRgridderinv2(empgrid=uv.grid)
plot(uv, pch=16, col=rgb(0,0,0,0.1), xlim=c(0,1), ylim=c(0,1),
     xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
lines(qua.regressCOP(f=0.5, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.2, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.7, cop=composite2COP, para=para), col=2)

```

```

lines(qua.regressCOP(f=0.1, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.9, cop=composite2COP, para=para), col=2)

med.wrtu <- EMPIRqua.regress(f=0.5, empinv=uv.inv1)
lines(med.wrtu, col=2, lwd=4)
qua.wrtu <- EMPIRqua.regress(f=0.2, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.7, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.1, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=4)
qua.wrtu <- EMPIRqua.regress(f=0.9, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=4)

lines(qua.regressCOP2(f=0.5, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.2, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.7, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.1, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.9, cop=composite2COP, para=para), col=4)

med.wrtv <- EMPIRqua.regress2(f=0.5, empinv=uv.inv2)
lines(med.wrtv, col=4, lwd=4)
qua.wrtv <- EMPIRqua.regress2(f=0.2, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.7, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.1, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=4)
qua.wrtv <- EMPIRqua.regress2(f=0.9, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=4)#
## End(Not run)

```

EMPIRgridderinv2

Derivative Inverses of the Grid of the Bivariate Empirical Copula for U with respect to V

Description

Generate a gridded representation of the inverse of the derivatives of the *bivariate empirical copula* of U with respect to V . This function is the empirical analog to [derCOPinv2](#).

Usage

```
EMPIRgridderinv2(empgrid=NULL, kumaraswamy=FALSE, dergrid=NULL, ...)
```

Arguments

empgrid	The grid from EMPIRgrid ;
kumaraswamy	A logical to trigger Kumaraswamy smoothing of the conditional quantile function;

dergrid	The results of EMPIRgridder2 and if left NULL then that function is called internally. There is some fragility at times in the quality of the numerical derivative and the author has provided this argument so that the derivative can be computed externally and then fed to this inversion function; and
...	Additional arguments to pass.

Value

The gridded values of the inverse of the derivative of U with respect to V .

Author(s)

W.H. Asquith

See Also

[EMPIRcop](#), [EMPIRcopdf](#), [EMPIRgrid](#), [EMPIRgridder2](#), [EMPIRgridderinv](#), [EMPIRgridderinv2](#)

Examples

```
# See examples under EMPIRgridderinv
```

EMPIRgrid_fast

Fast Grid of the Bivariate Empirical Copula

Description

Generate a fast gridded representation of the *bivariate empirical copula* (Hernández-Maldonado *et al.*, 2024). Mathematics for the "1/n" and "bernstein" copula are shown under [EMPIRcop](#). Herein, the $\ddot{C}_n$ (double-dots) are used to denote gridding of the copula. This function is a much more recent addition (spring 2026) to the **copBasic** than its historical ancestor of [EMPIRgrid](#) because of publication of the algorithm by Hernández-Maldonado *et al.* (2024). The **copBasic** author personally thanks V.M. Hernández-Maldonado for a May 2026 discussion on the algorithm and some clarifications to the pseudo-code in Hernández-Maldonado *et al.* (2024).

Usage

```
EMPIRgrid_fast(uv=NULL, ctype=c("1/n", "bernstein"), para=NULL,
               gridonly=FALSE, verbose=FALSE, ...)
```

Arguments

uv	A data frame or matrix of u and v vectors of nonexceedance probabilities (see also para);
ctype	A means to trigger which definition of empirical copula is to be used: $C_n(u, v)$ ("1/n") or $C_n^B(u, v)$ ("bernstein");

para	A matrix of u and v vectors as an alternative to uv for syntax compatibility for other situations in copBasic ;
verbose	A logical controlling whether the progress during grid building is to be shown;
gridonly	A logical to return the empcop component ($\check{C}_n(u, v)$ or $\check{C}_n^B(u, v)$); and
...	Additional arguments to pass to EMPIRcop .

Value

A matrix is returned ($\check{C}_n(u, v)$ or $\check{C}_n^B(u, v)$), if `gridonly` is true. Otherwise a list of the u , v , and $\check{C}_n(u, v)$ or $\check{C}_n^B(u, v)$ is returned. The list will also contain the `deluv` (simply $1/n$ for sample size n) used and the `ctype`. The list is returned as classes of "empirical.copula.grid" and "list". This is classification might be used in **copBasic** to check for a grid in other functions. Note, there is no `deluv` argument to this function like there is for the historical function **EMPIRgrid**.

Author(s)

W.H. Asquith

References

Hernández-Maldonado, V.M., Erdely, A., Díaz-Viera, M., and Rios, L., 2024, Fast procedure to compute empirical and Bernstein copulas: Applied Mathematics and Computation, v. 477, article 128827, 14 p., doi:10.1016/j.amc.2024.128827.

See Also

EMPIRgrid, **EMPIRgrid_lkup**

Examples

```
para <- list(cop=GHcop, para=3, breve=0.1)
uv <- simCOP(10, cop=breveCOP, para=para)
gf <- EMPIRgrid_fast(para=uv, gridonly=TRUE)
gg <- EMPIRgrid(para=uv, gridonly=TRUE, deluv=0.1, ctype="1/n")
print(sum(gf - gg)) # should be zero

n <- 5; uv <- matrix(runif(n * 2), ncol=2)
empCOP <- EMPIRgrid_fast(uv, ctype="1/n",          gridonly=TRUE)
          image(empCOP, main="1/n"                )
empCOP <- EMPIRgrid_fast(uv, ctype="bernstein", gridonly=TRUE)
          image(empCOP, main="Bernstein")

## Not run:
# This permutation asymmetric copula is shown as a means to cross check
# the orientation of the matrix computations within the function.
n <- 500; para <- list(cop=GHcop, para=c(12,15), breve=0.5)
uv <- copBasic::simCOP(n, cop=breveCOP, para=para, graphics=FALSE)
empCOP <- EMPIRgrid_fast(uv, gridonly=TRUE)
image(empCOP); points(uv) #
## End(Not run)
```

EMPIRgrid_lkup	<i>Empirical Copula by Bilinear Interpolation of Gridded Bivariate Empirical Copula</i>
----------------	---

Description

Compute joint probability through a gridded *empirical copula* $C_n(u, v)$ using bilinear interpolation (vectorized, no for looping). The function is designed to act like [EMPIRcop](#), but in lieu of consuming the bivariate data in `para` argument, the `EMPIRgrid_lkup` operates off a precomputed gridded list representation of the empirical copula computed either from [EMPIRgrid](#) (legacy) or [EMPIRgrid_fast](#). Given a gridded empirical copula denoted as $\ddot{C}_n$ and bounding cells of a rectangle 11 (lower left), 12 (lower right), 21 (upper left), and 22, the $C_n(u, v)$ can be estimated by bilinear interpolation in a weighted mean form as:

$$C_n(u, v) \approx w_{11} \times \ddot{C}_{11} + w_{12} \times \ddot{C}_{12} + w_{21} \times \ddot{C}_{21} + w_{22} \times \ddot{C}_{22},$$

for which the weights are

$$w_{11} = \frac{(u_2 - u)(v_2 - v)}{(u_2 - u_1)(v_2 - v_1)}, w_{12} = \frac{(u_2 - u)(v - v_1)}{(u_2 - u_1)(v_2 - v_1)},$$

$$w_{21} = \frac{(u - u_1)(v_2 - v)}{(u_2 - u_1)(v_2 - v_1)}, \text{ and } w_{22} = \frac{(u - u_1)(v - v_1)}{(u_2 - u_1)(v_2 - v_1)}.$$

Usage

```
EMPIRgrid_lkup(u, v, para=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A gridded empirical data structure from EMPIRgrid and EMPIRgrid_fast ; and
<code>...</code>	Additional arguments to pass to EMPIRgrid_fast if <code>para</code> is null.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

See Also

[EMPIRgrid](#), [EMPIRgrid_fast](#)

Examples

```
## Not run:
n <- 2000
para <- list(cop=GHcop, para=c(2.3,1.5), breve=0.5)
uv <- simCOP(n, cop=breveCOP, para=para, graphics=TRUE)
system.time(gridn <- EMPIRgrid(uv, ctype="1/n", deluv=1/2000))
system.time(gridf <- EMPIRgrid_fast(uv, ctype="1/n"))
# gridf should be shown to be over 100 times faster.
copn <- EMPIRgrid_lkup(uv[,1], uv[,2], para=gridn)
copf <- EMPIRgrid_lkup(uv[,1], uv[,2], para=gridf)
plot(copn, copf); abline(0,1) # equal value line
## End(Not run)

## Not run:
n <- 21 # Make a small sample demonstration and exhaustive list of
# comparisons in empirical copula computation styles.
para <- list(cop=GHcop, para=c(2.3,1.5), breve=0.5)
uv <- simCOP(n, cop=breveCOP, para=para, graphics=TRUE )
gridn <- EMPIRgrid_fast(uv, ctype="1/n", gridonly=FALSE)
gridw <- EMPIRgrid_fast(uv, ctype="bernstein", gridonly=FALSE)
copw <- EMPIRcop(uv[,1], uv[,2], para=uv, ctype="weibull" )
copn <- EMPIRcop(uv[,1], uv[,2], para=uv, ctype="1/n" )
coph <- EMPIRcop(uv[,1], uv[,2], para=uv, ctype="hazen" )
copb <- EMPIRcop(uv[,1], uv[,2], para=uv, ctype="bernstein" )
copc <- EMPIRcop(uv[,1], uv[,2], para=uv, ctype="checkerboard")
copgn <- EMPIRgrid_lkup(uv[,1], uv[,2], para=gridn)
copgb <- EMPIRgrid_lkup(uv[,1], uv[,2], para=gridw)

plot( copgn, copw, pch=24, col="seagreen4", bg="seagreen2")
points(copgn, copn, pch=24, col="red4", bg="red1" )
points(copgn, coph, pch=24, col="blue4", bg="blue1" )
points(copgn, copb, pch=24, col="orchid4", bg="orchid1" )
points(copgn, copc, pch=24, col="salmon4", bg="salmon1" )

points(copgb, copw, pch=25, col="seagreen", bg="white")
points(copgb, copn, pch=25, col="red", bg="white")
points(copgb, coph, pch=25, col="blue", bg="white")
points(copgb, copb, pch=25, col="orchid1", bg="white")
points(copgb, copc, pch=25, col="salmon", bg="white")

abline(0,1, lty=2) # equal value line
## End(Not run)
```

EMPIRmed.regress

*Median Regression of the Grid of the Bivariate Empirical Copula for
V with respect to U*

Description

Perform *median regression* from the gridded inversion of the *bivariate empirical copula* of V with respect to U .

Usage

```
EMPIRmed.regress(...)
```

Arguments

... Arguments to pass to [EMPIRqua.regress](#).

Value

The gridded values of the median regression of V with respect to U .

Author(s)

W.H. Asquith

See Also

[EMPIRgridderinv](#), [EMPIRqua.regress](#), [EMPIRmed.regress2](#), [EMPIRmed.regress2](#)

Examples

```
# See examples under EMPIRqua.regress
```

EMPIRmed.regress2	<i>Median Regression of the Grid of the Bivariate Empirical Copula for U with respect to V</i>
-------------------	--

Description

Perform *median regression* from the gridded inversion of the *bivariate empirical copula* of U with respect to V .

Usage

```
EMPIRmed.regress2(...)
```

Arguments

... Arguments to pass to [EMPIRqua.regress2](#).

Value

The gridded values of the median regression of U with respect to V .

Author(s)

W.H. Asquith

See Also

[EMPIRgridderv2](#), [EMPIRqua.regress](#), [EMPIRmed.regress](#), [EMPIRmed.regress2](#)

Examples

```
# See examples under EMPIRqua.regress
```

EMPIRqua.regress	<i>Quantile Regression of the Grid of the Bivariate Empirical Copula for V with respect to U</i>
------------------	--

Description

Perform *quantile regression* from the gridded inversion of the *bivariate empirical copula* of V with respect to U .

Usage

```
EMPIRqua.regress(f=0.5, u=seq(0.01,0.99, by=0.01), empinv=NULL,
                 lowess=FALSE, f.lowess=1/5, ...)
```

Arguments

<code>f</code>	The nonexceedance probability F to perform regression at and defaults to median regression $F = 1/2$;
<code>u</code>	A vector of u nonexceedance probabilities;
<code>empinv</code>	The grid from EMPIRgridderv2 ;
<code>lowess</code>	Perform lowess smooth on the quantile regression using the smooth factor of <code>f.lowess</code> ;
<code>f.lowess</code>	Smooth factor of almost the same argument name fed to the <code>lowess()</code> function in R; and
<code>...</code>	Additional arguments to pass.

Value

The gridded values of the quantile regression of V with respect to U .

Author(s)

W.H. Asquith

See Also

[EMPIRgridderv2](#), [EMPIRqua.regress2](#), [EMPIRmed.regress](#), [EMPIRmed.regress2](#)

Examples

```
## Not run: # EXAMPLE 1
theta <- 25
uv <- simCOP(n=1000, cop=PLACKETTcop, para=theta, ploton=FALSE, points=FALSE)
fakeU <- lmomco::pp(uv[,1], sort=FALSE)
fakeV <- lmomco::pp(uv[,2], sort=FALSE)
uv <- data.frame(U=fakeU, V=fakeV)

uv.grid <- EMPIRgrid(para=uv, deluv=0.05) # CPU hungry
uv.inv1 <- EMPIRgridderinv(empgrid=uv.grid)
plot(uv, pch=16, col=rgb(0,0,0,.1), xlim=c(0,1), ylim=c(0,1),
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
lines(qua.regressCOP(f=0.5, cop=PLACKETTcop, para=theta), lwd=2)
lines(qua.regressCOP(f=0.2, cop=PLACKETTcop, para=theta), lwd=2)
lines(qua.regressCOP(f=0.7, cop=PLACKETTcop, para=theta), lwd=2)
lines(qua.regressCOP(f=0.1, cop=PLACKETTcop, para=theta), lwd=2)
lines(qua.regressCOP(f=0.9, cop=PLACKETTcop, para=theta), lwd=2)

med.wrtu <- EMPIRqua.regress(f=0.5, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(med.wrtu, col=2, lwd=4)
qua.wrtu <- EMPIRqua.regress(f=0.2, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.7, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.1, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=4)
qua.wrtu <- EMPIRqua.regress(f=0.9, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=4)

library(quantreg) # Quantile Regression by quantreg
U <- seq(0.01, 0.99, by=0.01)
rq1m <- rq(V~U, data=uv, tau=0.1)
rq.1 <- rq1m$coefficients[1] + rq1m$coefficients[2]*U
rq1m <- rq(V~U, data=uv, tau=0.2)
rq.2 <- rq1m$coefficients[1] + rq1m$coefficients[2]*U
rq1m <- rq(V~U, data=uv, tau=0.5)
rq.5 <- rq1m$coefficients[1] + rq1m$coefficients[2]*U
rq1m <- rq(V~U, data=uv, tau=0.7)
rq.7 <- rq1m$coefficients[1] + rq1m$coefficients[2]*U
rq1m <- rq(V~U, data=uv, tau=0.9)
rq.9 <- rq1m$coefficients[1] + rq1m$coefficients[2]*U

lines(U, rq.1, col=4, lwd=2, lty=4)
lines(U, rq.2, col=4, lwd=2, lty=2)
lines(U, rq.5, col=4, lwd=4)
lines(U, rq.7, col=4, lwd=2, lty=2)
lines(U, rq.9, col=4, lwd=2, lty=4)#
## End(Not run)

## Not run: # EXAMPLE 2
# Start again with the PSP copula
uv <- simCOP(n=10000, cop=PSP, ploton=FALSE, points=FALSE)
```

```

fakeU <- lmomco::pp(uv[,1], sort=FALSE)
fakeV <- lmomco::pp(uv[,2], sort=FALSE)
uv <- data.frame(U=fakeU, V=fakeV)

uv.grid <- EMPIRgrid(para=uv, deluv=0.05) # CPU hungry
uv.inv1 <- EMPIRgridderinv(empgrid=uv.grid)
plot(uv, pch=16, col=rgb(0,0,0,0.1), xlim=c(0,1), ylim=c(0,1),
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
lines(qua.regressCOP(f=0.5, cop=PSP), lwd=2)
lines(qua.regressCOP(f=0.2, cop=PSP), lwd=2)
lines(qua.regressCOP(f=0.7, cop=PSP), lwd=2)
lines(qua.regressCOP(f=0.1, cop=PSP), lwd=2)
lines(qua.regressCOP(f=0.9, cop=PSP), lwd=2)

med.wrtu <- EMPIRqua.regress(f=0.5, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(med.wrtu, col=2, lwd=4)
qua.wrtu <- EMPIRqua.regress(f=0.2, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.7, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.1, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=4)
qua.wrtu <- EMPIRqua.regress(f=0.9, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=4)

library(quantreg) # Quantile Regression by quantreg
U <- seq(0.01, 0.99, by=0.01)
rq1m <- rq(V~U, data=uv, tau=0.1)
rq.1 <- rq1m$coefficients[1] + rq1m$coefficients[2]*U
rq1m <- rq(V~U, data=uv, tau=0.2)
rq.2 <- rq1m$coefficients[1] + rq1m$coefficients[2]*U
rq1m <- rq(V~U, data=uv, tau=0.5)
rq.5 <- rq1m$coefficients[1] + rq1m$coefficients[2]*U
rq1m <- rq(V~U, data=uv, tau=0.7)
rq.7 <- rq1m$coefficients[1] + rq1m$coefficients[2]*U
rq1m <- rq(V~U, data=uv, tau=0.9)
rq.9 <- rq1m$coefficients[1] + rq1m$coefficients[2]*U

lines(U, rq.1, col=4, lwd=2, lty=4)
lines(U, rq.2, col=4, lwd=2, lty=2)
lines(U, rq.5, col=4, lwd=4)
lines(U, rq.7, col=4, lwd=2, lty=2)
lines(U, rq.9, col=4, lwd=2, lty=4)#
## End(Not run)

## Not run: # EXAMPLE 3
uv <- simCOP(n=10000, cop=PSP, ploton=FALSE, points=FALSE)
fakeU <- lmomco::pp(uv[,1], sort=FALSE)
fakeV <- lmomco::pp(uv[,2], sort=FALSE)
uv <- data.frame(U=fakeU, V=fakeV)

uv.grid <- EMPIRgrid(para=uv, deluv=0.1) # CPU hungry
uv.inv1 <- EMPIRgridderinv(empgrid=uv.grid)

```

```

uv.inv2 <- EMPIRgridderinv2(empgrid=uv.grid)
plot(uv, pch=16, col=rgb(0,0,0,.1), xlim=c(0,1), ylim=c(0,1),
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
lines(qua.regressCOP(f=0.5, cop=PSP), col=2)
lines(qua.regressCOP(f=0.2, cop=PSP), col=2)
lines(qua.regressCOP(f=0.7, cop=PSP), col=2)
lines(qua.regressCOP(f=0.1, cop=PSP), col=2)
lines(qua.regressCOP(f=0.9, cop=PSP), col=2)

med.wrtu <- EMPIRqua.regress(f=0.5, empinv=uv.inv1)
lines(med.wrtu, col=2, lwd=4)
qua.wrtu <- EMPIRqua.regress(f=0.2, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.7, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.1, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=4)
qua.wrtu <- EMPIRqua.regress(f=0.9, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=4)

lines(qua.regressCOP2(f=0.5, cop=PSP), col=4)
lines(qua.regressCOP2(f=0.2, cop=PSP), col=4)
lines(qua.regressCOP2(f=0.7, cop=PSP), col=4)
lines(qua.regressCOP2(f=0.1, cop=PSP), col=4)
lines(qua.regressCOP2(f=0.9, cop=PSP), col=4)

med.wrtv <- EMPIRqua.regress2(f=0.5, empinv=uv.inv2)
lines(med.wrtv, col=4, lwd=4)
qua.wrtv <- EMPIRqua.regress2(f=0.2, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.7, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.1, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=4)
qua.wrtv <- EMPIRqua.regress2(f=0.9, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=4)#
## End(Not run)

## Not run: # EXAMPLE 4
# Now try a much more complex shape
# lowess smoothing on quantile regression is possible,
# see next example
para <- list(alpha=0.15, beta=0.65,
             cop1=PLACKETTcop, cop2=PLACKETTcop, para1=0.005, para2=1000)
uv <- simCOP(n=20000, cop=composite2COP, para=para)
fakeU <- lmomco::pp(uv[,1], sort=FALSE)
fakeV <- lmomco::pp(uv[,2], sort=FALSE)
uv <- data.frame(U=fakeU, V=fakeV)

uv.grid <- EMPIRgrid(para=uv, deluv=0.025) # CPU hungry
uv.inv1 <- EMPIRgridderinv(empgrid=uv.grid)
uv.inv2 <- EMPIRgridderinv2(empgrid=uv.grid)
plot(uv, pch=16, col=rgb(0,0,0,.1), xlim=c(0,1), ylim=c(0,1),

```

```

      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
lines(qua.regressCOP(f=0.5, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.2, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.7, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.1, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.9, cop=composite2COP, para=para), col=2)

med.wrtu <- EMPIRqua.regress(f=0.5, empinv=uv.inv1)
lines(med.wrtu, col=2, lwd=4)
qua.wrtu <- EMPIRqua.regress(f=0.2, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.7, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.1, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=4)
qua.wrtu <- EMPIRqua.regress(f=0.9, empinv=uv.inv1)
lines(qua.wrtu, col=2, lwd=2, lty=4)

lines(qua.regressCOP2(f=0.5, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.2, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.7, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.1, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.9, cop=composite2COP, para=para), col=4)

med.wrtv <- EMPIRqua.regress2(f=0.5, empinv=uv.inv2)
lines(med.wrtv, col=4, lwd=4)
qua.wrtv <- EMPIRqua.regress2(f=0.2, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.7, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.1, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=4)
qua.wrtv <- EMPIRqua.regress2(f=0.9, empinv=uv.inv2)
lines(qua.wrtv, col=4, lwd=2, lty=4)#
## End(Not run)

## Not run: # EXAMPLE 5
plot(uv, pch=16, col=rgb(0,0,0,.1), xlim=c(0,1), ylim=c(0,1),
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
lines(qua.regressCOP(f=0.5, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.2, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.7, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.1, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.9, cop=composite2COP, para=para), col=2)

med.wrtu <- EMPIRqua.regress(f=0.5, empinv=uv.inv1, lowess=TRUE)
lines(med.wrtu, col=2, lwd=4)
qua.wrtu <- EMPIRqua.regress(f=0.2, empinv=uv.inv1, lowess=TRUE)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.7, empinv=uv.inv1, lowess=TRUE)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.1, empinv=uv.inv1, lowess=TRUE)
lines(qua.wrtu, col=2, lwd=2, lty=4)

```

```

qua.wrtu <- EMPIRqua.regress(f=0.9, empinv=uv.inv1, lowess=TRUE)
lines(qua.wrtu, col=2, lwd=2, lty=4)

lines(qua.regressCOP2(f=0.5, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.2, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.7, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.1, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.9, cop=composite2COP, para=para), col=4)

med.wrtv <- EMPIRqua.regress2(f=0.5, empinv=uv.inv2, lowess=TRUE)
lines(med.wrtv, col=4, lwd=4)
qua.wrtv <- EMPIRqua.regress2(f=0.2, empinv=uv.inv2, lowess=TRUE)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.7, empinv=uv.inv2, lowess=TRUE)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.1, empinv=uv.inv2, lowess=TRUE)
lines(qua.wrtv, col=4, lwd=2, lty=4)
qua.wrtv <- EMPIRqua.regress2(f=0.9, empinv=uv.inv2, lowess=TRUE)
lines(qua.wrtv, col=4, lwd=2, lty=4)#
## End(Not run)

## Not run: # EXAMPLE 6 (changing the smoothing on the lowess)
plot(uv, pch=16, col=rgb(0,0,0,0.1), xlim=c(0,1), ylim=c(0,1),
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
lines(qua.regressCOP(f=0.5, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.2, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.7, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.1, cop=composite2COP, para=para), col=2)
lines(qua.regressCOP(f=0.9, cop=composite2COP, para=para), col=2)

med.wrtu <- EMPIRqua.regress(f=0.5, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(med.wrtu, col=2, lwd=4)
qua.wrtu <- EMPIRqua.regress(f=0.2, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.7, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=2)
qua.wrtu <- EMPIRqua.regress(f=0.1, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=4)
qua.wrtu <- EMPIRqua.regress(f=0.9, empinv=uv.inv1, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtu, col=2, lwd=2, lty=4)

lines(qua.regressCOP2(f=0.5, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.2, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.7, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.1, cop=composite2COP, para=para), col=4)
lines(qua.regressCOP2(f=0.9, cop=composite2COP, para=para), col=4)

med.wrtv <- EMPIRqua.regress2(f=0.5, empinv=uv.inv2, lowess=TRUE, f.lowess=0.1)
lines(med.wrtv, col=4, lwd=4)
qua.wrtv <- EMPIRqua.regress2(f=0.2, empinv=uv.inv2, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtv, col=4, lwd=2, lty=2)
qua.wrtv <- EMPIRqua.regress2(f=0.7, empinv=uv.inv2, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtv, col=4, lwd=2, lty=2)

```

```
qua.wrtv <- EMPIRqua.regress2(f=0.1, empinv=uv.inv2, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtv, col=4, lwd=2, lty=4)
qua.wrtv <- EMPIRqua.regress2(f=0.9, empinv=uv.inv2, lowess=TRUE, f.lowess=0.1)
lines(qua.wrtv, col=4, lwd=2, lty=4)#
## End(Not run)
```

EMPIRqua.regress2	<i>Quantile Regression of the Grid of the Bivariate Empirical Copula for U with respect to V</i>
-------------------	--

Description

Generate quantile regression from the gridded inversion of the *bivariate empirical copula* of U with respect to V .

Usage

```
EMPIRqua.regress2(f=0.5, v=seq(0.01,0.99, by=0.01), empinv=NULL,
                  lowess=FALSE, f.lowess=1/5, ...)
```

Arguments

<code>f</code>	The nonexceedance probability F to perform regression at and defaults to median regression $F = 1/2$;
<code>v</code>	A vector of v nonexceedance probabilities;
<code>empinv</code>	The grid from EMPIRgridderinv ;
<code>lowess</code>	Perform lowess smooth on the quantile regression using the smooth factor of <code>f=f.lowess</code> ;
<code>f.lowess</code>	Smooth factor of almost the same argument name fed to the <code>lowess()</code> function in R;
<code>...</code>	Additional arguments to pass.

Value

The gridded values of the quantile regression of U with respect to V .

Author(s)

W.H. Asquith

See Also

[EMPIRgridderinv2](#), [EMPIRqua.regress](#), [EMPIRmed.regress](#), [EMPIRmed.regress2](#)

Examples

```
# See examples under EMPIRqua.regress
```

Description

EXPERIMENTAL—Perform a simulation on a *bivariate empirical copula* to produce the random variates U and V and return an `R data.frame` of them. The method is more broadly known as *conditional simulation method*. This function is an empirical parallel to `simCOP` that is used for parametric copulas. If circumstances require conditional simulation of $V|U$, then function `EMPIRsimv`, which produces a vector of V from a fixed u , should be used.

For the usual situation in which an individual u during the simulation loops is not a value aligned on the grid, then the bounding conditional quantile functions are solved for each of the n simulations and the following interpolation is made by

$$v = \frac{v_1/w_1 + v_2/w_2}{1/w_1 + 1/w_2},$$

which states that the weighted mean is computed. The values v_1 and v_2 are ordinates of the conditional quantile function for the respective grid lines to the left and right of the u value. The values $w_1 = u - u_{\text{grid}}^{\text{left}}$ and $w_2 = u_{\text{grid}}^{\text{right}} - u$.

Usage

```
EMPIRsim(n=100, empgrid=NULL, kumaraswamy=FALSE, na.rm=TRUE, kept=FALSE,
         graphics=TRUE, ploton=TRUE, points=TRUE, snv=FALSE,
         infsnv.rm=TRUE, trapinfsnv=.Machine$double.eps, ...)
```

Arguments

<code>n</code>	A sample size, default is 100;
<code>empgrid</code>	Gridded empirical copula from <code>EMPIRgrid</code> ;
<code>kumaraswamy</code>	A logical to trigger Kumaraswamy distribution smoothing of the conditional quantile function that is passed to <code>EMPIRgridderinv</code> . The Kumaraswamy distribution is a distribution having support $[0, 1]$ with an explicit quantile function and takes the place of a Beta distribution (see lmomco function <code>quakur()</code> for more details);
<code>na.rm</code>	A logical to toggle the removal of NA entries on the returned <code>data.frame</code> ;
<code>kept</code>	Keep the t uniform random variable for the simulation as the last column in the returned <code>data.frame</code> ;
<code>graphics</code>	A logical that will disable graphics by setting <code>ploton</code> and <code>points</code> to <code>FALSE</code> and overriding whatever their settings were;
<code>ploton</code>	A logical to toggle on the plot;
<code>points</code>	A logical to actually draw the simulations by the <code>points()</code> function in <code>R</code> ;

<code>snv</code>	A logical to convert the $\{u, v\}$ to standard normal scores (variates) both for the optional graphics and the returned <code>data.frame</code> . Joe (2014) advocates extensively for use of normal scores, which is in contrast to Nelsen (2006) who does not;
<code>infsnv.rm</code>	A logical that will quietly strip out any occurrences of $u = \{0, 1\}$ or $v = \{0, 1\}$ from the simulations because these are infinity in magnitude when converted to standard normal variates is to occur. Thus, this logical only impacts logic flow when <code>snv</code> is TRUE. The <code>infsnv.rm</code> is mutually exclusive from <code>trapinfsnv</code> ;
<code>trapinfsnv</code>	If TRUE and presumably small, the numerical value of this argument (η) is used to replace $u = \{0, 1\}$ and $v = \{0, 1\}$ with $u(0) = v(0) = \eta$ or $u(1) = v(1) = 1 - \eta$ as appropriate when conversion to standard normal variates is to occur. The setting of <code>trapinfsnv</code> only is used if <code>snv</code> is TRUE and <code>infsnv.rm</code> is FALSE; and
<code>...</code>	Additional arguments to pass to the <code>points()</code> function in R.

Value

An R `data.frame` of the simulated values is returned.

Author(s)

W.H. Asquith

See Also

[EMPIRgrid](#), [EMPIRgridderinv](#), [EMPIRsimv](#)

Examples

```
# See other examples under EMPIRsimv

## Not run:
pdf("EMPIRsim_experiment.pdf")
nsim <- 5000
para <- list(alpha=0.15, beta=0.65,
             cop1=PLACKETTcop, cop2=PLACKETTcop, para1=0.005, para2=1000)
set.seed(1)
uv <- simCOP(n=nsim, cop=composite2COP, para=para, snv=TRUE,
            pch=16, col=rgb(0,0,0,.2))
mtext("A highly complex simulated bivariate relation")
# set.seed(1) # try not resetting the seed
uv.grid <- EMPIRgrid(para=uv, deluv=0.025)

uv2 <- EMPIRsim(n=nsim, empgrid=uv.grid, kumaraswamy=FALSE, snv=TRUE,
               col=rgb(1,0,0,0.1), pch=16)
mtext("Resimulation without Kumaraswamy smoothing")

uv3 <- EMPIRsim(n=nsim, empgrid=uv.grid, kumaraswamy=TRUE, snv=TRUE,
               col=rgb(1,0,0,0.1), pch=16)
mtext("Resimulation but using the Kumaraswamy Distribution for smoothing")
```

```
dev.off()#
## End(Not run)
```

EMPIRsimv

Simulate a Bivariate Empirical Copula For a Fixed Value of U

Description

EXPERIMENTAL—Perform a simulation on a *bivariate empirical copula* to extract the random variates V from a given and fixed value for $u = \text{constant}$. The purpose of this function is to return a simple vector of the V simulations. This behavior is similar to [simCOPmicro](#) but differs from the general 2-D simulation implemented in the other functions: [EMPIRsim](#) and [simCOP](#)—these two functions generate R data.frames of simulated random variates U and V and optional graphics as well.

For the usual situation in which u is not a value aligned on the grid, then the bounding conditional quantile functions are solved for each of the n simulations and the following interpolation is made by

$$v = \frac{v_1/w_1 + v_2/w_2}{1/w_1 + 1/w_2},$$

which states that the weighted mean is computed. The values v_1 and v_2 are ordinates of the conditional quantile function for the respective grid lines to the left and right of the u value. The values $w_1 = u - u_{\text{grid}}^{\text{left}}$ and $w_2 = u_{\text{grid}}^{\text{right}} - u$.

Usage

```
EMPIRsimv(u, n=1, empgrid=NULL, kumaraswamy=FALSE, ...)
```

Arguments

<code>u</code>	The fixed probability u on which to perform conditional simulation for a sample of size n ;
<code>n</code>	A sample size, default is 1;
<code>empgrid</code>	Gridded empirical copula from EMPIRgrid ;
<code>kumaraswamy</code>	A logical to trigger Kumaraswamy distribution smoothing of the conditional quantile function that is passed to EMPIRgridderinv . The Kumaraswamy distribution is a distribution having support $[0, 1]$ with an explicit quantile function and takes the place of a Beta distribution (see lmomco function <code>quakur()</code> for more details); and
<code>...</code>	Additional arguments to pass.

Value

A vector of simulated V values is returned.

Author(s)

W.H. Asquith

See Also

[EMPIRgrid](#), [EMPIRsim](#)

Examples

```
## Not run:
nsim <- 3000
para <- list(alpha=0.15, beta=0.65,
             cop1=PLACKETTcop, cop2=PLACKETTcop, para1=.005, para2=1000)
set.seed(10)
uv <- simCOP(n=nsim, cop=composite2COP, para=para, pch=16, col=rgb(0,0,0,0.2))
uv.grid <- EMPIRgrid(para=uv, deluv=.1)
set.seed(1)
V1 <- EMPIRsimv(u=0.6, n=nsim, empgrid=uv.grid)
set.seed(1)
V2 <- EMPIRsimv(u=0.6, n=nsim, empgrid=uv.grid, kumaraswamy=TRUE)
plot(V1,V2)
abline(0,1)

invgrid1 <- EMPIRgridderinv(empgrid=uv.grid)
invgrid2 <- EMPIRgridderinv(empgrid=uv.grid, kumaraswamy=TRUE)
att <- attributes(invgrid2); kur <- att$kumaraswamy
# Now draw random variates from the Kumaraswamy distribution using
# rlmomco() and vec2par() provided by the lmomco package.
set.seed(1)
kurpar <- lmomco::vec2par(c(kur$Alpha[7], kur$Beta[7]), type="kur")
Vsim <- lmomco::rlmomco(nsim, kurpar)

print(summary(V1)) # Kumaraswamy not core in QDF reconstruction
print(summary(V2)) # Kumaraswamy core in QDF reconstruction
print(summary(Vsim)) # Kumaraswamy use of the kumaraswamy

# Continuing with a conditional quantile 0.74 that will not land along one of the
# grid lines, a weighted interpolation will be done.
set.seed(1) # try not resetting the seed
nsim <- 5000
V <- EMPIRsimv(u=0.74, n=nsim, empgrid=uv.grid)
# It is important that the uv.grid used to make V is the same grid used in inversion
# with kumaraswamy=TRUE to guarantee that the correct Kumaraswamy parameters are
# available if a user is doing cut and paste and exploring these examples.
set.seed(1)
V1 <- lmomco::rlmomco(nsim, lmomco::vec2par(c(kur$Alpha[8], kur$Beta[8]), type="kur"))
set.seed(1)
V2 <- lmomco::rlmomco(nsim, lmomco::vec2par(c(kur$Alpha[9], kur$Beta[9]), type="kur"))

plot( lmomco::pp(V), sort(V), type="l", lwd=4, col=8) # GREY is empirical from grid
lines(lmomco::pp(V1), sort(V1), col=2, lwd=2) # Kumaraswamy at u=0.7 # RED
lines(lmomco::pp(V2), sort(V2), col=3, lwd=2) # Kumaraswamy at u=0.8 # GREEN

W1 <- 0.74 - 0.7; W2 <- 0.8 - 0.74
Vblend <- (V1/W1 + V2/W2) / sum(1/W1 + 1/W2)
lines(lmomco::pp(Vblend), sort(Vblend), col=4, lwd=2) # BLUE LINE
```

```
# Notice how the grey line and the blue diverge for about F < 0.1 and F > 0.9.
# These are the limits of the grid spacing and linear interpolation within the
# grid intervals is being used and not direct simulation from the Kumaraswamy.
## End(Not run)
```

EuvCOP

Expected value of U given V

Description

Compute the *expected value* of U given a V (the Y direction) through the *conditional distribution function* $G(Y)$ using the appropriate *partial derivative* of a copula ($\mathbf{C}(u, v)$) with respect to V . The inversion of the partial derivative is the *conditional quantile function*. Basic principles provide the expectation for a $y \geq 0$ is

$$E[Y] = \int_0^\infty y f(y) dy = \int_0^\infty (1 - G_y(Y)) dy,$$

which for the setting here becomes

$$E[U \mid V = v] = \int_0^1 \left(1 - \frac{\delta}{\delta v} \mathbf{C}(u, v)\right) du.$$

This function solves the integral using the [derCOP2](#) function. This avoids a call of the [derCOPinv2](#) through its `uniroot()` inversion of the derivative. The example shown for `EuvCOP()` below does a validation check using conditional simulation, which is dependence (of course) of the design of the **copBasic** package, as part of simple isolation of a horizontal slice of the simulation and computing the mean of the V within the slice.

Usage

```
EuvCOP(v=seq(0.01, 0.99, by=0.01), cop=NULL, para=NULL, asuv=FALSE, nsim=1E5,
        subdivisions=200L, rel.tol=.Machine$double.eps^0.25, abs.tol=rel.tol, ...)
```

Arguments

<code>v</code>	Nonexceedance probability v in the Y direction;
<code>cop</code>	A copula function with vectorization as in <code>asCOP</code> ;
<code>para</code>	Vector of parameters or other data structures, if needed, to pass to the copula;
<code>asuv</code>	Return a data frame of the U and V ;
<code>nsim</code>	Number of simulations for <i>Monte Carlo integration</i> if numerical integration were to fail (refer to Note);
<code>subdivisions</code>	Argument of same name passed to <code>integrate()</code> ;
<code>rel.tol</code>	Argument of same name passed to <code>integrate()</code> ;
<code>abs.tol</code>	Argument of same name passed to <code>integrate()</code> ; and
<code>...</code>	Additional arguments to pass to derCOP2 .

Value

Value(s) for the expectation are returned.

Note

The author is well aware that the name of this function does not contain the number 2 as the family of functions also sharing this *with respect to v* nature. It was a design mistake in 2008 to have used the 2. The uv in the function name is the moniker for this *with respect to v*.

There can be the rare examples of the numerical integration failing. In such circumstances, Monte Carlo integration is performed and the returned vector becomes a named vector with the `sim` identifying values stemming from the simulation.

```
para <- list(cop=RFcop, para=0.9)
para <- list(cop=COP, para=para, reflect=1, alpha=0, beta=0.3)
EuvCOP(c(0.0001, 0.0002, 0.001, 0.01, 0.1), cop=composite1COP, para=para)
#           sim
#[1] 0.001319395 0.002238238 0.006905300 0.034608078 0.173451788
```

Author(s)

W.H. Asquith

See Also

[EuvCOP](#), [derCOP2](#)

Examples

```
# We can show algorithmic validation using a highly asymmetric case of a
# copula having its parameter also nearly generating a singular component.
v <- c(0.2, 0.8); n <- 5E2; set.seed(1)
para <- list(cop=HRcop, para=120, alpha=0.4, beta=0.05)
UV <- simCOP(n, cop=composite1COP, para=para, graphics=FALSE) # set TRUE to view

sapply(v, function(vv) EuvCOP(vv, cop=composite1COP, para=para))
# [1] 0.3051985 0.7059999

sapply(v, function(vv) mean( UV$U[UV$V > vv - 50/n & UV$V < vv + 50/n] ) )
# [1] 0.2796518 0.7092755 # general validation is thus shown as n-->large

# If visualized, we see in the lower corner than heuristics suggest a mean further
# to the right of the "singularity" for v=0.2 than v=0.80. For v=0.80, the
# "singularity" appears tighter given the upper tail dependency contrast of the
# coupla in the symmetrical case (alpha=0, beta=0) and changing the parameter to
# a Spearman Rho (say) similar to the para setting in this example. So, 0.70 for
# the mean given v=0.80 makes sense. Further notice that the two estimates of the
# mean are further apparent for v=0.2 relative to v=0.80. Again, this makes sense
# when the copula is visualized even at small n let alone large n.

# See additional Examples under EuvCOP().
```

```
## Not run:
set.seed(1)
n <- 5000; Vlo <- rep(0.001, n); Vhi <- rep(0.95, n); Theta <- 3
Ulo <- simCOPmicro(Vlo, cop=J0copB5, para=Theta); dlo <- density(Ulo)
Uhi <- simCOPmicro(Vhi, cop=J0copB5, para=Theta); dhi <- density(Uhi)
dlo$x[dlo$x < 0] <- 0; dhi$x[dhi$x < 0] <- 0
dlo$x[dlo$x > 1] <- 1; dhi$x[dhi$x > 1] <- 1

summary(Ulo)
Ulo_mu <- EuvCOP(Vlo[1], cop=J0copB5, para=Theta); print(Ulo_mu)
#      Min.    1st Qu.      Median        Mean     3rd Qu.        Max.
# 0.0000669 0.0887330 0.2006123 0.2504796 0.3802847 0.9589315
#
#      Ulo_mu -----> 0.2502145
summary(Uhi)
Uhi_mu <- EuvCOP(Vhi[1], cop=J0copB5, para=Theta); print(Uhi_mu)
#      Min.    1st Qu.      Median        Mean     3rd Qu.        Max.
# 0.01399 0.90603 0.93919 0.9157600 0.95946 0.99411
#
#      Uhi_mu -----> 0.9154093

UV <- simCOP(n, cop=J0copB5, para=Theta,
             cex=0.6, pch=21, bg="palegreen", col="darkgreen")
abline(h=Vlo[1], col="salmon", lty=3) # near the bottom to form datum for density
abline(h=Vhi[1], col="purple", lty=3) # near the top to form datum for density
lines(dlo$x, dlo$y/max(dlo$y)/2 + Vlo[1], col="salmon", lwd=2)
# re-scaled density along the line already drawn near the bottom (Vlo)
# think rug plotting to bottom the values plotting very close to the line
lines(dhi$x, 1-dhi$y/max(dhi$y)/2 - (1-Vhi[1]), col="purple", lwd=2)
# re-scaled density along the line already drawn near the top (Vhi)
# think rug plotting to bottom the values plotting very close to the line
uv <- seq(0.001, 0.999, by=0.001) # for trajectory of E[U | V=v]
lines(EuvCOP(uv, cop=J0copB5, para=Theta), uv, col="blue", lwd=3.5, lty=2)
points(Ulo_mu, Vlo[1], pch=16, col="salmon", cex=2)
points(Uhi_mu, Vhi[1], pch=16, col="purple", cex=2) #
## End(Not run)
```

EuvCOP

Expected value of V given U

Description

Compute the *expected value* of V given a U (the X direction) through the *conditional distribution function* $F(X)$ using the appropriate *partial derivative* of a copula ($C(u, v)$) with respect to U . The inversion of the partial derivative is the *conditional quantile function*. Basic principles provide the expectation for a $x \geq 0$ is

$$E[X] = \int_0^\infty x f(x) dx = \int_0^\infty (1 - F_x(X)) dx,$$

which for the setting here becomes

$$E[V \mid U = u] = \int_0^1 \left(1 - \frac{\delta}{\delta u} C(u, v)\right) dv.$$

This function solves the integral using the [derCOP](#) function. Verification study is provided in the **Note** section.

Usage

```
EvuCOP(u=seq(0.01, 0.99, by=0.01), cop=NULL, para=NULL, asuv=FALSE, nsim=1E5,
        subdivisions=200L, rel.tol=.Machine$double.eps^0.25, abs.tol=rel.tol, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>cop</code>	A copula function with vectorization as in <code>asCOP</code> ;
<code>para</code>	Vector of parameters or other data structures, if needed, to pass to the copula;
<code>asuv</code>	Return a data frame of the U and V ;
<code>nsim</code>	Number of simulations for <i>Monte Carlo integration</i> if numerical integration were to fail (refer to Note in EvuCOP);
<code>subdivisions</code>	Argument of same name passed to <code>integrate()</code> ;
<code>rel.tol</code>	Argument of same name passed to <code>integrate()</code> ;
<code>abs.tol</code>	Argument of same name passed to <code>integrate()</code> ; and
<code>...</code>	Additional arguments to pass to derCOP .

Value

Value(s) for the expectation are returned.

Note

For the [PSP](#) copula with no parameters, compute the the median and mean V given $U = 0.4$, respectively:

```
U <- 0.4; n <- 1E4
med.regressCOP(u=U, cop=PSP) # V = 0.4912752
set.seed(1)
median(replicate(n, derCOPinv(cop=PSP, U, runif(1)) ) ) # V = 0.4876440
mean( replicate(n, derCOPinv(cop=PSP, U, runif(1)) ) ) # V = 0.5049729
```

It is seen in the above that the median V given U is very close to the mean, but is not equal. Using the derivative inversion within [med.regressCOP](#) the median is about 0.491 and then using large-sample simulation, about 0.491 too is computed. This confirms the median and long standing proven use of [derCOP](#) (conditional distribution function) and [derCOPinv](#) (conditional quantile function) within the package. The expectation (mean) by simulation provides the anchor point to check implementation of `EuvCOP()`. The mean for V given U is about 0.505. Continuing, the core logic of `EvuCOP()` is to use numerical integration of the conditional distribution function (the partial derivative) and not bother for speed purposes to use the inversion of the partial derivative:

```

integrate(function(v)                                1-derCOP(  cop=PSP, U, v),
          lower=0, upper=1) # 0.5047805 with absolute error < 1.4e-11

integrate(function(v) sapply(v, function(t) derCOPinv(cop=PSP, U, t)),
          lower=0, upper=1) # 0.5047862 with absolute error < 7.2e-05

```

The two integrals match, which functions as a confirmation of the $(1 - F)$ term in the mathematical definition. Finally, the two integrals match the simulation results. The expectation or mean $V | U = 0.4$ for the [PSP](#) copula is about 0.5048.

Author(s)

W.H. Asquith

See Also

[EuvCOP](#), [derCOP](#)

Examples

```

# Highly asymmetric and reflected Clayton copula for which visualization
para <- list(cop=CLcop, para=30, alpha=0.2, beta=0.6, reflect=3)
# UV <- simCOP(5000, cop=breveCOP, para=para, cex=0.5); abline(v=0.25, col="red")
EuvCOP(0.25, cop=breveCOP, para=para) # 0.5982261
# confirms that at U=0.25 that an intuitive estimate would be about 0.6.

## Not run:
# Secondary validation of the EuvCOP() and EuvCOP() implementation
UV <- simCOP(200, cop=PSP)
u <- seq(0.005, 0.995, by=0.005)
v <- sapply(u, function(t) integrate(function(k)
                                     1 - derCOP(cop=PSP, t, k), lower=0, upper=1)$value)
lines(u,v, col="red", lwd=7, lty=1) # red line
v <- seq(0.005, 0.995, by=0.005)
u <- sapply(v, function(t) integrate(function(k)
                                     1 - derCOP2(cop=PSP, k, t), lower=0, upper=1)$value)
lines(u,v, col="red", lwd=7, lty=2) # dashed red line

uv <- seq(0.005, 0.995, by=0.005) # solid and dashed white lines
lines(EvuCOP(uv, cop=PSP, asuv=TRUE), col="white", lwd=3, lty=1)
lines(EuvCOP(uv, cop=PSP, asuv=TRUE), col="white", lwd=3, lty=3)

# median regression lines for comparison, green and green dashed lines
lines(med.regressCOP( uv, cop=PSP), col="seagreen", lwd=1.5, lty=1)
lines(med.regressCOP2(uv, cop=PSP), col="seagreen", lwd=1.5, lty=4) #
## End(Not run)

## Not run:
uv <- seq(0.005, 0.995, by=0.005) # stress testing eample with singularity
UV <- simCOP(50, cop=M_N5p12b, para=2)
lines(EvuCOP(uv, cop=M_N5p12b, para=2, asuv=TRUE), col="red", lwd=5)
lines(EuvCOP(uv, cop=M_N5p12b, para=2, asuv=TRUE), col="skyblue", lwd=1) #

```

```
## End(Not run)

## Not run:
uv <- seq(0.005, 0.995, by=0.005) # more asymmetry ---- mean regression in UV and VU
para <- list(cop=GHcop, para=23, alpha=0.1, beta=0.6, reflect=3)
para <- list(cop=breveCOP, para=para)
UV <- simCOP(200, cop=COP, para=para)
lines(EuvCOP(uv, cop=COP, para=para, asuv=TRUE), col="red", lwd=2)
lines(EvuCOP(uv, cop=COP, para=para, asuv=TRUE), col="blue", lwd=2) #
## End(Not run)

## Not run:
# Open questions? The derCOP() and derCOPinv() functions of the package have long
# been known to work "properly." But let us think again on the situation of
# permutation symmetry about the equal value line. Recalling that this symmetry is
# orthogonal to the equal value line, it remains open whether there could be
# asymmetry in the vertical (or horizontal). Let us draw some median regression lines
# and see that they do not plot perfectly on the equal value line, but could this be
# down to numerical issues and by association the simulation of the copula itself
# that is also using derCOPinv() (conditional simulation method). Then, we can plot
# the expectations and we see that these are not equal to the medians, but again are
# close. *** Do results here indicate edges of numerical performance? ***
t <- seq(0.01, 0.99, by=0.01)
UV <- simCOP(10000, cop=N4212cop, para=4, pch=21, lwd=0.8, col=8, bg="white")
lines(med.regressCOP(cop=N4212cop, para=4, asuv=TRUE), col="red")
lines(med.regressCOP2(cop=N4212cop, para=4, asuv=TRUE), col="red")
abline(0, 1, col="deepskyblue", lwd=3); abline(v=0.5, col="deepskyblue", lwd=4)
lines(EvuCOP(t, cop=N4212cop, para=4, asuv=TRUE), pch=16, col="darkgreen")
lines(EuvCOP(t, cop=N4212cop, para=4, asuv=TRUE), pch=16, col="darkgreen") #
## End(Not run)
```

FGMcop

The Generalized Farlie–Gumbel–Morgenstern Copula

Description

The *generalized Farlie–Gumbel–Morgenstern copula* (Bekrizade *et al.*, 2012) is

$$\mathbf{C}_{\Theta, \alpha, n}(u, v) = \mathbf{FGM}(u, v) = uv \left[1 + \Theta(1 - u^\alpha)(1 - v^\alpha) \right]^n,$$

where $\Theta \in [-\min\{1, 1/(n\alpha^2)\}, +1/(n\alpha)]$, $\alpha > 0$, and $n \in 0, 1, 2, \dots$. The copula $\Theta = 0$ or $\alpha = 0$ or $n = 0$ becomes the *independence copula* ($\Pi(u, v)$; **P**). When $\alpha = n = 1$, then the well-known, single-parameter Farlie–Gumbel–Morgenstern copula results, and *Spearman Rho* (**rhoCOP**) is $\rho_C = \Theta/3$ but in general

$$\rho_C = 12 \sum_{r=1}^n \binom{n}{r} \Theta^r \left[\frac{\Gamma(r+1)\Gamma(2/\alpha)}{\alpha\Gamma(r+1+2/\alpha)} \right]^2.$$

The support of $\rho_C(\cdot \cdot \cdot ; \Theta, 1, 1)$ is $[-1/3, +1/3]$ but extends via α and n to $\approx [-0.50, +0.43]$, which shows that the generalization of the copula increases the range of dependency. The generalized version is implemented by FGMcop.

The *iterated Farlie–Gumbel–Morgenstern copula* (Chine and Benatia, 2017) for the r th iteration is

$$C_{\beta}(u, v) = \mathbf{FGMi}(u, v) = uv + \sum_{j=1}^r \beta_j \cdot (uv)^{[j/2]} \cdot (u'v')^{[(j+1)/2]},$$

where $u' = 1 - u$ and $v' = 1 - v$ for $|\beta_j| \leq 1$ that has r dimensions $\beta = (\beta_1, \dots, \beta_j, \dots, \beta_r)$ and $[t]$ is the integer part of t . The copula $\beta = 0$ becomes the *independence copula* ($\mathbf{\Pi}(u, v)$; **P**). The support of $\rho_{\mathbf{C}}(\dots; \beta)$ is approximately $[-0.43, +0.43]$. The iterated version is implemented by FGMicop. Internally, the r is determined from the length of the β in the para argument.

Usage

```
FGMcop( u, v, para=c(NA, 1,1), ...)
FGMicop(u, v, para=NULL,      ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector of parameters. For the generalized version, the Θ , α , and n of the copula where the default argument shows the need to include the Θ . However, if a fourth parameter is present, it is treated as a logical to reverse the copula ($u + v - 1 + \mathbf{FGM}(1 - u, 1 - v; \Theta, \alpha, n)$). Also if a single parameter is given, then the $\alpha = n = 1$ are automatically set to produce the single-parameter Farlie–Gumbel–Morgenstern copula. For the iterated version, the β vector of r iterations;
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Bekrizade, Hakim, Parham, G.A., Zadkarmi, M.R., 2012, The new generalization of Farlie–Gumbel–Morgenstern copulas: *Applied Mathematical Sciences*, v. 6, no. 71, pp. 3527–3533.

Chine, Amel, and Benatia, Fatah, 2017, Bivariate copulas parameters estimation using the trimmed L-moments methods: *Afrika Statistika*, v. 12, no. 1, pp. 1185–1197, [doi:10.16929/as/2017.1185.99](https://doi.org/10.16929/as/2017.1185.99).

See Also

[P](#), [mleCOP](#)

Examples

```
## Not run:
# Bekrizade et al. (2012, table 1) report for a=2 and n=3 that range in
# theta = [-0.1667, 0.1667] and range in rho = [-0.1806641, 0.4036458]. However,
# we see that they have seemingly made an error in listing the lower bounds of theta:
rhoCOP(FGMcop, para=c( 1/6, 2, 3)) # 0.4036458
rhoCOP(FGMcop, para=c( -1/6, 2, 3)) # Following error results
# In cop(u, v, para = para, ...) : parameter Theta < -0.0833333333333333
rhoCOP(FGMcop, para=c(-1/12, 2, 3)) # -0.1806641
## End(Not run)

## Not run:
# Support of FGMrcop(): first for r=1 iterations and then for large r.
sapply(c(-1, 1), function(t) rhoCOP(cop=FGMicop, para=rep(t, 1)) )
# [1] -0.3333333 0.3333333
sapply(c(-1, 1), function(t) rhoCOP(cop=FGMicop, para=rep(t,50)) )
# [1] -0.4341385 0.4341385
## End(Not run)

## Not run:
# Maximum likelihood estimation near theta upper bounds for a=3 and n=2.
set.seed(832)
UV <- simCOP(3000, cop=FGMcop, para=c(+0.16, 3, 2))
# Define a transform function for parameter domain, though mleCOP does
# provide some robustness anyway---not forcing n into the positive
# domain via as.integer(exp(p[3])) seems to not always be needed.
FGMpfunc <- function(p) {
  d <- p[1]; a <- exp(p[2]); n <- as.integer(exp(p[3]))
  lwr <- -min(c(1,1/(n*a^2))); upr <- 1/(n*a)
  d <- ifelse(d <= lwr, lwr, ifelse(d >= upr, upr, d))
  return( c(d, a, n) )
}
para <- c(0.16, 3, 2); init <- c(0, 1, 1)
ML <- mleCOP(UV$U, UV$V, cop=FGMcop, init.para=init, parafn=FGMpfunc)
print(ML$para) # [1] 0.1596361 3.1321228 2.0000000
# So, we have recovered reasonable estimates of the three parameters
# given through MLE estimation.
densityCOPplot(cop=FGMcop, para= para, contour.col="red")
densityCOPplot(cop=FGMcop, para=ML$para, ploton=FALSE) #
## End(Not run)

## Not run:
# L-comoment ratio diagram space of Generalized Farlie-Gumbel-Morgenstern Copula
FGM <- NULL
for(n in seq(1, 100, by=1)) { Fgm <- NULL # to support intermediate plotting
  for(a in 10^seq(-5, 5, by=0.02)) { if(a == 0) next # in case log10 not involved
    rng <- c(-pmin(1, 1/(n*a^2)), 1/(n*a))
    ts <- runif(10, min=rng[1], max=rng[2])
    for(t in ts) {
      fgm <- lcomCOP(cop=FGMcop, para=c(t, a, n))
      fgm <- data.frame(t, a, n, T2.12=fgm$lcomUV[2], T2.21=fgm$lcomVU[2],
        T3.12=fgm$lcomUV[3], T3.21=fgm$lcomVU[3],
```

```

                                T4.12=fgm$lcomUV[4], T4.21=fgm$lcomVU[4])
    row.names(fgm) <- NULL; Fgm <- rbind(Fgm, fgm)
  }
}
plot(c(-0.5, +0.5), c(-0.1, 0.20), type="n", las=1, main=n,
      xlab="Spearman Rho", ylab="L-coskew and L-cokurtosis")
points(Fgm$T2.12, Fgm$T3.12, col="seagreen3")
points(Fgm$T2.12, Fgm$T4.12, col="orchid3" )
FGM <- rbind(FGM, Fgm)
}
plot(c(-0.5, +0.5), c(-0.1,0.20), type="n", las=1,
      xlab="Spearman Rho", ylab="L-coskew and L-cokurtosis")
points(FGM$T2.12, FGM$T3.12, col="seagreen3")
points(FGM$T2.12, FGM$T4.12, col="orchid3" )
save(FGM, file="LCM_FGM.RData") #
## End(Not run)

```

footCOP

The Spearman Footrule of a Copula

Description

Compute the measure of association known as the *Spearman Footrule* $\psi_{\mathbf{C}}$ (Nelsen *et al.*, 2001, p. 281), which is defined as

$$\psi_{\mathbf{C}} = \frac{3}{2} \mathcal{Q}(\mathbf{C}, \mathbf{M}) - \frac{1}{2},$$

where $\mathbf{C}(u, v)$ is the copula, $\mathbf{M}(u, v)$ is the *Fréchet–Hoeffding upper bound* (**M**), $\mathcal{Q}(a, b)$ is a *concordance function* (**concordCOP**) (Nelsen, 2006, p. 158), and $-0.5 \leq \psi_{\mathbf{C}} \leq 1$. The $\psi_{\mathbf{C}}$ in terms of a single integration pass on the copula (Genest *et al.*, 2010, p. 940) is

$$\psi_{\mathbf{C}} = 1 - \int_{\mathcal{T}^2} |u - v| d\mathbf{C}(u, v) = 6 \int_0^1 \mathbf{C}(u, u) du - 2.$$

Note, Nelsen *et al.* (2001) use $\phi_{\mathbf{C}}$ but that symbol is taken in **copBasic** for the *Hoeffding Phi* (**hoefCOP**), and Spearman Footrule does not seem to appear in Nelsen (2006). From the definition, Spearman Footrule only depends on the *primary diagonal* (alt. *main diagonal*, Genest *et al.*, 2010) of the copula, $\mathbf{C}(t, t)$ (**diagCOP**). The function here supports the first $\psi_{\mathbf{C}}$ computation by `as.concordance=TRUE`, but the default is the later definition by the direct integration on the diagonal.

Usage

```
footCOP(cop=NULL, para=NULL, by.concordance=FALSE, as.sample=FALSE, ...)
```

Arguments

<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>by.concordance</code>	Instead of using the single integral to compute ψ_C , use the concordance function method implemented through concordCOP ; and
<code>as.sample</code>	A logical controlling whether an optional R data.frame in <code>para</code> is used to compute the $\hat{\psi}$ (see Note); and
<code>...</code>	Additional arguments to pass, which are dispatched to the copula function <code>cop</code> and possibly concordCOP , such as <code>brute</code> or <code>delta</code> used by that function.

Value

The value for ψ_C is returned.

Note

Conceptually, the sample Spearman Footrule is a standardized sum of the absolute differences in the ranks (Genest *et al.*, 2010, eq. 1). The sample $\hat{\psi}$ is

$$\hat{\psi} = 1 - \frac{3}{n^2 - 1} \sum_{i=1}^n |R_i - S_i|,$$

where R_i and S_i are the respective ranks of X and Y , n is sample size, and $-0.5 \lesssim \hat{\psi} \leq 1$. Genest *et al.* (2010, p. 937) also remark that other normalizations have been used in the literature. The approximate $\hat{\psi}$ lower bounds can be seen by the following where -0.5 is achieved always for odd sample sizes and rapidly approached as sample size grows:

```
sams <- seq_len(25) # samples from 1 to 25 in size
minFoot <- sapply(sams, function(i) 1 - (3/(i^2-1))*sum(abs((1:i)-(i:1))))
plot(sams, minFoot, type="b", col="brown4")
lines(par()$usr[1:2], rep(-0.5, 2), lty=2)
```

The sample $\hat{\psi}$ also can be written (Genest *et al.*, 2010, eq. 3) as

$$\hat{\psi} = \frac{6}{n-1} \left[\sum_i^n \min\left(\frac{R_i}{n+1}, \frac{S_i}{n+1}\right) \right] - \frac{2n+1}{n-1},$$

which is not implemented here because the first sample version is modestly faster. Also, the sampling variance of $\hat{\psi}$ under *assumption of independence* between X and Y ($E[\hat{\psi}] = 0$) is

$$\text{var}(\hat{\psi}) = \frac{2n^2 + 7}{5(n+1)(n-1)^2}.$$

Genest *et al.* (2010, p. 938) say that prior literature shows that in small samples, Spearman Footrule is less variable than the well-known *Spearman Rho* ([rhoCOP](#)). For a copula having continuous partial derivatives, then as $n \rightarrow \infty$, the quantity $(\hat{\psi} - \psi_C)\sqrt{n} \sim \text{Normal}(0, \text{var}(\gamma_C))$. Genest *et al.* (2010)

show variance of $\hat{\psi}$ for the *independence copula* ($C(u, v) = \Pi(u, v)$, **P**) as $\text{var}(\psi_C) = 2/5$. For comparison, the *Gini Gamma* for independence is larger at $\text{var}(\gamma_C) = 2/3$ (see [giniCOP Note](#)). Genest *et al.* (2010) also present additional material for estimation of the distribution $\hat{\psi}$ variance for conditions of dependence based on copulas. In Genest *et al.* (2010, p. 941), independence and two examples of dependence, $\text{var}(\hat{\gamma}) > \text{var}(\hat{\psi})$, but those authors do not appear to remark on whether this inequality holds for all copula.

Author(s)

W.H. Asquith

References

- Beliakov, G., de Amo, E., Juan Fernández-Sánchez, J., Úbeda-Flores, M., 2022, Best-possible bounds on the set of copulas with a given value of Spearman's footrule, *Fuzzy Sets and Systems*, v. 428, pp. 138–152, [doi:10.1016/j.fss.2020.11.011](https://doi.org/10.1016/j.fss.2020.11.011).
- Genest, C., Nešlehová, J., and Ghorbal, N.B., 2010, Spearman's footrule and Gini's gamma—A review with complements: *Journal of Nonparametric Statistics*, v. 22, no. 8, pp. 937–954, [doi:10.1080/10485250903499667](https://doi.org/10.1080/10485250903499667).
- Nelsen, R.B., 2006, *An introduction to copulas*: New York, Springer, 269 p.
- Nelsen, R.B., Quesada-Molina, J.J., Rodríguez-Lallena, J.A., Úbeda-Flores, M., 2001, Distribution functions of copulas—A class of bivariate probability integral transforms: *Statistics and Probability Letters*, v. 54, no. 3, pp. 277–282, [doi:10.1016/S01677152\(01\)000608](https://doi.org/10.1016/S01677152(01)000608).

See Also

[blomCOP](#), [giniCOP](#), [hoefCOP](#), [rhoCOP](#), [tauCOP](#), [wolfCOP](#)

Examples

```
footCOP(cop=PSP) # 0.3177662
# footCOP(cop=PSP, by.concordance=TRUE) # 0.3178025 # slower, likely giving up accuracy

## Not run:
set.seed(1); n <- 2000; UV <- simCOP(n=n, cop=GHCop, para=2.3, graphics=FALSE)
footCOP(para=UV, as.sample=TRUE) # 0.5356089 (sample version)
footCOP(cop=GHCop, para=2.3) # 0.5513380 (copula integration)
footCOP(cop=GHCop, para=2.3, by.concordance=TRUE) # 0.5513562 (concordance function)
# where the later issued warnings on the integration
## End(Not run)

## Not run:
# Sweep through the Plackett as means to show -0.5 <= Footrule <= 1, and to show
for(p in seq(-4,4, by=0.5)) { # that the sample version is in agreement.
  p <- round(10^p, digits=4); UV <- simCOP(10000, cop=PLcop, para=p, lwd=0.5)
  footS <- round(footCOP(as.sample=TRUE, para=UV), digits=4)
  footT <- round(footCOP(cop=PLcop, para=p), digits=4)
  mtext(paste0("Plackett(theta=", p, ")"), line=1)
  mtext(paste0("Spearman Footrule : theoretical=", footT, ", sample=", footS))
} #
```

```
## End(Not run)

## Not run:
set.seed(1); nsim <- 1000
varFTunderIndepend <- function(n) {
  (2*n^2 + 7) / (5*(n+1)*(n-1)^2) # Genest et al. (2010)
}
ns <- c(10, 15, 20, 25, 50, 75, 100)
plot(min(ns):max(ns), varFTunderIndepend(10:max(ns)), type="l",
      xlab="Sample size", ylab="Variance of Sample Estimator", col="salmon4")
mtext("Sample Spearman Footrule Under Independence", col="salmon4")
for(n in ns) {
  sFT <- vector(length=nsim)
  for(i in seq_len(nsim)) {
    uv <- simCOP(n=n, cop=P, para=2, graphics=FALSE)
    sFT[i] <- footCOP(para=uv, as.sample=TRUE)
  }
  varFT <- varFTunderIndepend(n)
  zz <- round(c(n, mean(sFT), var(sFT), varFT), digits=6)
  names(zz) <- c("n", "mean_sim", "var_sim", "var_Genest")
  print(zz)
  points(n, zz[3], cex=2, pch=21, col="salmon4", bg="salmon1")
} # results show proper implementation and Genest et al. (2010, sec. 3)
## End(Not run)
```

Description

The *Frank copula* (Joe, 2014, p. 165) is

$$\mathbf{C}_{\Theta}(u, v) = \mathbf{FR}(u, v) = -\frac{1}{\Theta} \log \left[\frac{1 - e^{-\Theta} - (1 - e^{-\Theta u})(1 - e^{-\Theta v})}{1 - e^{-\Theta}} \right],$$

where $\Theta \in [-\infty, +\infty]$, $\Theta \neq 0$. The copula, as $\Theta \rightarrow -\infty$ limits, to the *countermonotonicity coupla* ($\mathbf{W}(u, v)$; **W**), as $\Theta \rightarrow 0^{\pm}$ limits to the *independence copula* ($\mathbf{\Pi}(u, v)$; **P**), and as $\Theta \rightarrow +\infty$, limits to the *comonotonicity copula* ($\mathbf{M}(u, v)$; **M**). The parameter Θ is readily computed from a *Kendall Tau* (**tauCOP**) by numerical methods as $\tau_{\mathbf{C}}(\Theta) = 1 + 4\Theta^{-1}[D_1(\Theta) - 1]$ or from a *Spearman Rho* (**rhoCOP**) as $\rho_{\mathbf{C}}(\Theta) = 1 + 4\Theta^{-1}[D_2(\Theta) - D_1(\Theta)]$ for *Debye function* as

$$D_k(x, k) = kx^{-k} \int_0^x t^k (e^t - 1)^{-1} dt.$$

Usage

```
FRcop(u, v, para=NULL, rhotau=NULL, userhotau_chk=TRUE,
      cortype=c("kendall", "spearman", "tau", "rho"), ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula;
<code>rhotau</code>	Optional Kendall Tau or Spearman Rho and parameter <code>para</code> is returned depending on the setting of <code>cortype</code> . The <code>u</code> and <code>v</code> can be used for estimation of the parameter as computed through the setting of <code>cortype</code> ;
<code>cortype</code>	A character string controlling, if the parameter is not given, to use a Kendall Tau or Spearman Rho for estimation of the parameter. The name of this argument is reflective of an internal call to <code>stats::cor()</code> to the correlation (association) setting for Kendall Tau or Spearman Rho;
<code>userhotau_chk</code>	A logical to trigger computation of Kendall Tau for the given parameter and used as a secondary check on numerical limits of the copula implementation for the package; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned. Otherwise if `tau` is given, then the Θ is computed and a list having

<code>para</code>	The parameter Θ , and
<code>tau</code>	Kendall Tau.

and if `para=NULL` and `tau=NULL`, then the values within `u` and `v` are used to compute Kendall Tau and then compute the parameter, and these are returned in the aforementioned list. Or if `rho` is given, then the Θ is computed and a similar list is returned having similar structure but with Spearman Rho instead.

Note

Whether because of default directions of derivative in [derCOP](#) for partial derivative of the copula or other numerical challenges, the implementation uses the negative parameter whether positive or not and rotates the copula as needed for complete operation of [simCOP](#). Several default limiting conditions are in the sources before conversion to independence, perfect positive dependence, or perfect negative dependence.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[M](#), [P](#), [W](#)

Examples

```

UV <- simCOP(n=1000, cop=FRcop, para=20, seed=1)
print(FRcop(UV[,1], UV[,2], cortype="kendall" )$tau) # 0.8072993
print(FRcop(UV[,1], UV[,2], cortype="spearman")$rho) # 0.9536648

## Not run:
# Joe (2014) example follows by extending the functionality of copBasic
# into a 3-dimensional C-vine copula using Frank and Gumbel copulas and
# Kendall Taus and Kendall Partial Taus. The example is expanded to show
# how advanced features of copBasic can be incorporated for asymmetry
# (if needed) and reflections (rotations) of copula:
nsim <- 5000 # number of simulations but not too large for long CPU
d <- 3       # three dimensions in Joe (2014, algorithm 15) implementation
GHcop_para <- GHcop(tau = 0.5, )$para # theta = 2
FRcop_para <- FRcop(rhotau=0.7, cortype="kendall")$para # theta = 11.41155
# Substantial notation complexity by structuring the C-vine by matrices for
# copula families and their parameters then dial in asymmetry by "breves"
# (see copBasic::breveCOP) and then reflection (rotation ability in the
# copulas themselves). With zero breves, we have no asymmetry (permutation)
# and we are going with the basic copula formula, so Reflects are all 1.
Cops <- matrix(c(NA, "GHcop", "FRcop",
                 NA, NA, "M",
                 NA, NA, NA), nrow=d, ncol=d, byrow=TRUE)
Thetas <- matrix(c(NA, GHcop_para, FRcop_para,
                  NA, NA, NA,
                  NA, NA, NA), nrow=d, ncol=d, byrow=TRUE)
Breves <- matrix(c(NA, 0, 0,
                  NA, NA, 0,
                  NA, NA, NA), nrow=d, ncol=d, byrow=TRUE)
Reflects <- matrix(c(NA, 1, 1,
                   NA, NA, 1,
                   NA, NA, NA), nrow=d, ncol=d, byrow=TRUE)
# see copBasic::breveCOP(), copBasic::GHcop(), copBasic::M()
# see also copBasic::COP() for how reflection work within the package
set.seed(1); U <- NULL # seed and vector of the U_{(1,2,3)} probabilities
for(i in 1:nsim) {
  w <- runif(d); u <- rep(NA, d); u[1] <- w[1]
  # looks messy but just a way dump a host of "parameters" including family
  # itself down into copBasic logic
  para <- list(cop=breveCOP, para=list(cop=eval(parse(text=Cops[1,2])),
                                     para=Thetas[1,2], breve=Breves[1,2], reflect=Reflects[1,2]))
  u[2] <- derCOPinv(u[1], t=w[2], cop=COP, para=para) # conditional quantiles
  for(j in 3:d) {
    q <- w[j]
    for(l in (j-1):1) { # Joe (2014, algorithm 16)
      para <- list(cop=breveCOP, para=list(cop=eval(parse(text=Cops[l,j])),
                                           para=Thetas[l,j], breve=Breves[l,j], reflect=Reflects[l,j]))
      q <- derCOPinv(w[l], t=q, cop=COP, para=para) # conditional quantiles
    }
    u[j] <- q
  }
  U <- rbind(U, matrix(u, ncol=d, byrow=TRUE))
}

```

```

}
U <- as.data.frame( U )
names( U ) <- paste0("U", seq_len(d))

# Kendall Partial Tau; Joe (2014, p. 8.66, Theorem 8.66)
"pc3dCOP" <- function(taujk, tauhj, tauhk) { # close attention to h subscript!
  etajk <- sin( pi*taujk / 2) # Expansion to trig by Joe (2014, theorem 8.19),
  etahj <- sin( pi*tauhj / 2) # Joe says this definition "might be"
  etahk <- sin( pi*tauhk / 2) # better than partial tau on scores.
  (2/pi) * asin( (etajk - etahj*etahk) / sqrt( (1-etahj^2) * (1-etahk^2) ) )
}
# Joe (2014, pp. 404-405)
# Kendall partial tau but needs pairwise Kendall taus to compute partial tau.
"PairwiseTaus" <- function(U) {
  ktau <- matrix(NA, nrow=d, ncol=d)
  for(j in 1:d) { for(l in 1:d) {
    if(j < l) next
    if(j == l) { ktau[l,j] <- 1; next } # 1s on diagonal as required.
    ktau[l,j] <- ktau[j,l] <- cor(U[,l], U[,j], method="kendall") # symmetrical
  } }
  return(ktau)
}
PairwiseTaus <- PairwiseTaus(U) # [1,2] and [1,3] are 0.5 and 0.7 matching
# requirement and more importantly for Joe (2014, table 8.3, p. 405). Now, [2,3] is
# about 0.783 which again matches Joe's results of 0.782.
Tau23given1pc <- pc3dCOP(PairwiseTaus[2,3], PairwiseTaus[2,1], PairwiseTaus[3,1])
# Joe (2014, table 8.3, p. 405) reports tau^{pc}_{jk|h} as 0.848 by giant
# simulation and we get about 0.852199 for some modest nsim and set.seed(1).
## End(Not run)

```

Description

The *Fréchet Family copula* (Durante, 2007, pp. 256–259) is

$$\mathbf{C}_{\alpha,\beta}(u,v) = \mathbf{FF}(u,v) = \alpha \mathbf{M}(u,v) + (1 - \alpha - \beta) \mathbf{\Pi}(u,v) + \beta \mathbf{W}(u,v),$$

where $\alpha, \beta \geq 0$ and $\alpha + \beta \leq 1$. The Fréchet Family copulas are *convex combinations* of the fundamental copulas $\mathbf{W}$ (*Fréchet–Hoeffding lower-bound copula*; **W**), $\mathbf{\Pi}$ (independence; **P**), and $\mathbf{M}$ (*Fréchet–Hoeffding upper-bound copula*; **M**). The copula is *comprehensive* because both $\mathbf{W}$ and $\mathbf{M}$ can be obtained. The parameters are readily estimated using *Spearman Rho* ($\rho_{\mathbf{C}}$; **rhoCOP**) and *Kendall Tau* ($\tau_{\mathbf{C}}$; **tauCOP**) by

$$\tau_{\mathbf{C}} = \frac{(\alpha - \beta)(\alpha + \beta + 2)}{3} \text{ and } \rho_{\mathbf{C}} = \alpha - \beta.$$

The Fréchet Family copula virtually always has a visible *singular component* unless $\alpha, \beta = 0$. The copula has respective *lower- and upper-tail dependency parameters* of $\lambda^L = \alpha$ and $\lambda^U = \alpha$ (**taildepCOP**). Durante (2007, p. 257) reports that the Fréchet Family copula can approximate any bivariate copula in a “unique way” and the error bound can be estimated.

Usage

```
FRECHETcop(u,v, para=NULL, rho=NULL, tau=NULL, par2rhotau=FALSE, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (two element) of parameters α and β ;
<code>rho</code>	Spearman Rho from which to estimate the parameters;
<code>tau</code>	Kendall Tau from which to estimate the parameters;
<code>par2rhotau</code>	A logical that if TRUE will return an R list of the ρ_C and τ_C for the parameters; and
<code>...</code>	Additional arguments to pass.

Details

The function will check the consistency of the parameters whether given by argument or computed from ρ_C and τ_C . The term “Family” is used with this particular copula in **copBasic** so as to draw distinction to the Fréchet lower- and upper-bound copulas as the two limiting copulas are called.

For no other reason than that it can be easily done and makes a nice picture, loop through a nest of ρ and τ for the Fréchet Family copula and plot the domain of the resulting parameters:

```
ops <- options(warn=-1) # warning supression because "loops" are dumb
taus <- rhos <- seq(-1,1, by=0.01)
plot(NA, NA, type="n", xlim=c(0,1), ylim=c(0,1),
     xlab="Frechet Copula Parameter Alpha",
     ylab="Frechet Copula Parameter Beta")
for(tau in taus) {
  for(rho in rhos) {
    fcop <- FRECHETcop(rho=rho, tau=tau)
    if(! is.na(fcop$para[1])) points(fcop$para[1], fcop$para[2])
  }
}
options(ops)
```

Value

Value(s) for the copula are returned using the α and β as set by argument `para`; however, if `para=NULL` and `rho` and `tau` are set and compatible with the copula, then $\{\rho_C, \tau_C\} \rightarrow \{\alpha, \beta\}$, parameter estimation made, and an R list is returned.

Note

A convex combination ([convex2COP](#)) of Π and $\mathbf{M}$, which is a modification of the Fréchet Family, is the *Linear Spearman* copula:

$$\mathbf{C}_\alpha(u, v) = (1 - \alpha)\Pi(u, v) + \alpha\mathbf{M}(u, v),$$

for $0 \leq \alpha \leq 1$, and the parameter is equal to ρ_C . When the convex combination is used for construction, the complement of the parameter is equal to ρ_C (e.g. $1 - \alpha = \rho_C$; `rhoCOP`), which can be validated by

```
rhoCOP(cop=convex2COP, para=list(alpha=1-0.48, cop1=P, cop2=M)) # 0.4799948
```

Author(s)

W.H. Asquith

References

Durante, F., 2007, Families of copulas, Appendix C, *in* Salvadori, G., De Michele, C., Kotegoda, N.T., and Rosso, R., 2007, *Extremes in Nature—An approach using copulas*: Springer, 289 p.

See Also

[M](#), [P](#), [W](#)

Examples

```
## Not run:
ppara <- c(0.25, 0.50)
fcop <- FRECHETcop(para=ppara, par2rho=TRUE)
RHO <- fcop$rho; TAU <- fcop$tau

level.curvesCOP(cop=FRECHETcop, para=ppara) # Durante (2007, Fig. C.27(b))
mtext("Frechet Family copula")
UV <- simCOP(n=50, cop=FRECHETcop, para=ppara, ploton=FALSE, points=FALSE)
tau <- cor(UV$U, UV$V, method="kendall" ) # sample Kendall Tau
rho <- cor(UV$U, UV$V, method="spearman") # sample Spearman Rho
spara <- FRECHETcop(rho=rho, tau=tau) # a fitted Frechet Family copula
spara <- spara$para
if(is.na(spara[1])) { # now a fittable combination is not guaranteed
  warning("sample rho and tau do not provide valid parameters, ",
    "try another simulation")
} else { # now if fit, draw some red-colored level curves for comparison
  level.curvesCOP(cop=FRECHETcop, para=spara, ploton=FALSE, col=2)
} #
## End(Not run)
```

Description

The *g-EV copula* (Joe, 2014, p. 105) is a limiting form of the *Normal (Gaussian) copula* ([NORMcop](#)):

$$C_\rho(u, v) = \mathbf{gEV}(u, v; \rho) = \exp(-A(x, y; \rho)),$$

where $x = -\log(u)$, $y = -\log(v)$, and

$$A(x, y; \rho) = y,$$

for $0 \leq x/(x+y) \leq \rho^2/(1+\rho^2)$,

$$A(x, y; \rho) = (x + y - 2\rho\sqrt{xy})/(1 - \rho^2),$$

for $\rho^2/(1+\rho^2) \leq x/(x+y) \leq 1/(1+\rho^2)$,

$$A(x, y; \rho) = x,$$

for $1/(1+\rho^2) \leq x/(x+y) \leq 1$ and where $\rho \in [0, 1]$. A somewhat curious observation is that this copula has relative hard boundaries into the upper-left and lower-right corners when compared to the other copulas supported by the **copBasic** package. In other words, the hull defined by the copula has a near hard (not fuzzy) curvilinear boundaries that adjust with the parameter ρ . The lower-case *g* in the copula name is deliberately to avoid the sequence GEV as to never be confused with the *generalized extreme value distribution*.

Usage

```
gEVcop(u, v, para=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	The parameter ρ ; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[tEVcop](#), [NORMcop](#)

Examples

```
## Not run:
UV <- simCOP(200, cop=gEVcop, para=0.8) #
## End(Not run)

## Not run:
# Joe (2014, p. 105) has brief detail indicating rho = [0,1] and though it seems
# Rho would be a Pearson correlation, this does not seem to be the case. The Rho
# seems to start with that of the Gaussian and then through the extreme-value
# transform, it just assumes the role of a parameter called Rho.
rho <- 0.8
UV <- simCOP(2000, cop=gEVcop, para=rho)
P <- cor(UV[,1], UV[,2], method="pearson")
if(abs(P - rho) < 0.001) {
  print("Yes same")
} else { print("nope not") } #
## End(Not run)

## Not run:
gEVparameter <- seq(0.02, 1, by=0.02)
SpearmanRho <- sapply(gEVparameter, function(k) rhoCOP(cop=gEVcop, para=k))
plot(gEVparameter, SpearmanRho)

for(rho in gEVparameter) {
  UV <- simCOP(n=1000, cop=gEVcop, para=rho)
  mtext(paste0("Parameter = ", rho))
} #
## End(Not run)
```

GHcop

The Gumbel–Hougaard Extreme Value Copula

Description

SYMMETRIC GUMBEL-HOUGAARD—The *Gumbel–Hougaard copula* (Nelsen, 2006, pp. 118 and 164) is

$$\mathbf{C}_{\Theta}(u, v) = \mathbf{GH}(u, v) = \exp \left\{ - \left[(-\log u)^{\Theta} + (-\log v)^{\Theta} \right]^{1/\Theta} \right\},$$

where $\Theta \in [1, \infty)$. The **GH** copula is a *bivariate extreme value copula (BEV)*. The parameter Θ is readily estimated using a *Kendall Tau* (say a sample version $\hat{\tau}$) where the τ of the copula ($\tau_{\mathbf{C}}$) is defined as

$$\tau_{\mathbf{C}} = \frac{\Theta - 1}{\Theta} \rightarrow \Theta = \frac{1}{1 - \tau}.$$

The copula is readily extended into d dimensions by

$$\mathbf{C}_{\Theta}(u, v) = \exp \left\{ - \left[(-\log u_1)^{\Theta} + \cdots + (-\log u_d)^{\Theta} \right]^{1/\Theta} \right\}.$$

However, such an implementation is not available in the **copBasic** package.

Every Gumbel–Hougaard copula is a *multivariate extreme value (MEV)* copula, and hence useful in analysis of extreme value distributions. The Gumbel–Hougaard copula is the *only* Archimedean *MEV* (Salvadori *et al.*, 2007, p. 192). The Gumbel–Hougaard copula has respective *lower-* and *upper-tail dependency* parameters of $\lambda^L = 0$ and $\lambda^U = 2 - 2^{1/\Theta}$, respectively. Nelsen (2006, p. 96) shows that $\mathbf{C}_\theta^r(u^{1/r}, v^{1/r}) = \mathbf{C}_\theta(u, v)$ so that every Gumbel–Hougaard copula has a property known as *max-stable*. A *dependence measure* uniquely defined for *BEV* copulas is shown under [rhobevCOP](#).

A comparison through simulation between Gumbel–Hougaard implementations by the R packages **acopula**, **copBasic**, **copula**, and **Gumbel** is shown in the **Examples** section. At least three divergent techniques for random variate generation are used amongst those packages. The simulations also use **copBasic**-style random variate generation (conditional simulation) using an analytical-numerical hybrid solution to conditional inverse described in the **Note** section.

TWO-PARAMETER GUMBEL–HOUGAARD—A *permutation symmetric* ([isCOP.permsym](#)) but almost certainly *radial asymmetric* ([isCOP.radsym](#)) version of the copula is readily constructed (Brahimi *et al.*, 2015) into a two-parameter version:

$$\mathbf{C}(u, v; \beta_1, \beta_2) = \left[\left((u^{-\beta_2} - 1)^{\beta_1} + (v^{-\beta_2} - 1)^{\beta_1} \right)^{1/\beta_1} + 1 \right]^{-1/\beta_2},$$

where $\beta_1 \geq 1$ and $\beta_2 > 0$. Both parameters controls the general level of association, whereas parameter β_2 can be thought of as controlling left-tail dependency ([taildepCOP](#), $\lambda_{(\beta_1, \beta_2)}^{[U|L]}$; e.g. $\lambda_{(1.5; \beta_2)}^U = 0.413$ for all β_2 but $\lambda_{(1.5; 0.2)}^L = 0.811$ and $\lambda_{(1.5; 2.2)}^L = 0.099$. Brahimi *et al.* (2015) report a *Spearman Rho* ([rhoCOP](#)) for a **GH**_(1.5, 0.2)(u, v) is 0.5, which is readily confirmed in **copBasic** by the function call `rhoCOP(cop=GHcop, para=c(1.5, 0.2))`. The two-parameter **GH** is triggered if the length of the `para` argument is exactly 2.

ASYMMETRIC GUMBEL–HOUGAARD—An asymmetric version of the copula is readily constructed (Joe, 2014, p. 185–186) into a three-parameter version with *Marshall–Olkin* copulas ([M0cop](#)) on the boundaries:

$$\mathbf{C}(u, v; \Theta, \pi_2, \pi_3) = \exp[-\mathcal{A}(-\log u, -\log v; \Theta, \pi_2, \pi_3)],$$

where $\Theta \geq 1$ as before, $0 \leq \pi_2, \pi_3 \leq 1$, and

$$\mathcal{A}(x, y; \Theta, \pi_2, \pi_3) = [(\pi_2 x)^\Theta + (\pi_3 y)^\Theta]^{1/\Theta} + (1 - \pi_2)x + (1 - \pi_3)y.$$

The asymmetric **GH** is triggered if the length of the `para` argument is exactly 3. The **GHcop** function provides no mechanism for estimation of the parameters for the asymmetric version. Reviewing simulations, the bounds on the π parameters in Joe (2014, p. 185) “[$0 \leq \pi_2 < \pi_3 \leq 1$]” might be incorrect—by Joe back referencing to Joe (2014, eq. 4.35, p. 183) the π -limits as stated for **copBasic** are shown. An algorithm for parameter estimation for the asymmetric **GH** using two different measures of *bivariate skewness* as well as an arbitrary *measure of association* is shown in section **Details** in [joeskewCOP](#).

Usage

```
GHcop(u, v, para=NULL, tau=NULL, tau.big=0.985, cor=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element or triplet) of parameters—the Θ parameter of the copula;
<code>tau</code>	Kendall Tau τ from which to estimate the parameter Θ ;
<code>tau.big</code>	The largest value for τ_C prior to switching to the M copula applicable to the symmetric version of this copula;
<code>cor</code>	A copBasic syntax for “the correlation coefficient” suitable for the copula—a synonym for <code>tau</code> ; and
<code>...</code>	Additional arguments to pass.

Details

Numerical experiments seem to indicate for $\tau_C > 0.985$ that failures in the numerical partial derivatives in `derCOP` and `derCOP2` result—a τ_C this large is indeed *large*. As $\Theta \rightarrow \infty$ the Gumbel–Hougaard copula becomes the *Fréchet–Hoeffding upper-bound copula* **M** (see [M](#)). A $\tau_C \approx 0.985$ yields $\Theta \approx 66 + 2/3$, then for $\Theta > 1/(1 - \tau_C)$ flips over to the **M** copula with a warning issued.

Value

Value(s) for the copula are returned using the Θ as set by argument `para`. Alternative returned values are possible: (1) If `para=NULL` and `tau` is set, then $\tau_C \rightarrow \Theta$ and an **R** list is returned. (2) If `para=NULL` and `tau=NULL`, then an attempt to estimate Θ from the u and v is made by $\text{cor}(u, v)_\tau \rightarrow \tau_C \rightarrow \Theta$ by either trigger using `cor(u, v, method="kendall")` in **R**, and an **R** list is returned. The possibly returned list has the following elements:

<code>para</code>	The computed Θ from the given bivariate data in <code>para</code> ; and
<code>tau</code>	The sample estimate of τ .

Note

SYMMETRIC GUMBEL–HOUGAARD — A function for the derivative of the copula (Joe, 2014, p. 172) given u is

```
"GHcop.derCOP" <- function(u, v, para=NULL, ...) {
  x <- -log(u); y <- -log(v)
  A <- exp(-(x^para + y^para)^(1/para)) * (1 + (y/x)^para)^(1/para - 1)
  return(A/u)
}
```

that can be tested by the following

```
Theta <- 1 / (1 - 0.15) # a Kendall Tau of 0.15
GHcop.derCOP( 0.5, 0.75, para=Theta) # 0.7787597
derCOP(cop=GHcop, 0.5, 0.75, para=Theta) # 0.7787597
# The next two nearly return same value but conversion to GRVs
```

```
# (Gumbel Reduced Variates) to magnify the numerical differences.
# The GHcop.derCOP is expected to be the more accurate of the two.
lmomco::prob2grv(GHcop.derCOP(      0.5, 0.9999999, para=Theta)) # 18.83349
lmomco::prob2grv(derCOP(cop=GHcop, 0.5, 0.9999999, para=Theta)) # 18.71497
lmomco::prob2grv(derCOP(cop=GHcop, 0.5, 0.9999999, para=Theta,
                        delu=.Machine$double.eps^.25)) # 18.83341
```

where the last numerical approximation shows that tighter tolerance is needed. A function for the inverse of the derivative (Joe, 2014, p. 172) given u by an analytical-numerical hybrid is

```
"GHcop.derCOPinv" <- function(u,t, para=NULL, verbose=FALSE,
                             tol=.Machine$double.eps, ...) {
  if(length(u) > 1) warning("only the first value of u will be used")
  if(length(t) > 1) warning("only the first value of t will be used")
  if(is.null(para)) { warning("para can not be NULL"); return(NA) }
  u <- u[1]; t <- t[1]; rt <- NULL
  x <- -log(u); A <- (x + (para - 1)*log(x) - log(t))
  hz <- function(z) { z + (para - 1)*log(z) - A }
  zmax <- x; i <- 0; hofz.lo <- hz(zmax)
  if(sign(hofz.lo) != -1) warning("sign for h(z) is not negative!")
  while(1) {
    i <- i + 1
    if(i > 100) {
      warning("maximum iterations looking for zmax reached"); break
    }
    # increment zmax by 1/2 log cycle, sign(hofz.lo) should be negative!
    if(sign(hz(zmax <- zmax + 1/2)) != sign(hofz.lo)) break
  }
  try(rt <- uniroot(hz, c(x, zmax), tol=tol, ...), silent=FALSE)
  if(verbose) print(rt)
  if(is.null(rt)) {
    warning("NULL on the inversion of the GH copula derivative")
    return(NA)
  }
  zo <- rt$root
  y <- (zo^para - x^para)^(1/para)
  names(y) <- NULL
  return(exp(-y))
}
```

that can be tested by the following, which also shows how to increase the tolerance on the numerical implementation

```
u <- 0.999; p <- 0.999
GHcop.derCOPinv(      u, p, para=1.56) # 0.999977
derCOPinv(cop=GHcop, u, p, para=1.56) # 1 (unity), needs tighter tolerance
derCOPinv(cop=GHcop, u, p, para=1.56, tol=.Machine$double.eps/10) # 0.999977
```

ASYMMETRIC GUMBEL-HOUGAARD — Set $\tau_C = 0.35$ then for a symmetric and then reflection

on the 1:1 line of the asymmetric Gumbel–Hougaard copula and compute the primary parameter Θ , and lastly, compute three bivariate ν_C skewnesses (`nuskewCOP`):

```
Theta1 <- uniroot(function(t) {
  0.35 - tauCOP(cop=GHcop, para=c(t)) }, c(1,10))$root
Theta2 <- uniroot(function(t) { # asymmetric
  0.35 - tauCOP(cop=GHcop, para=c(t, 0.6, 0.9)) }, c(1,30))$root
Theta3 <- uniroot(function(t) { # asymmetric reflection on 1:1
  0.35 - tauCOP(cop=GHcop, para=c(t, 0.9, 0.6)) }, c(1,30))$root
# Theta1 = 1.538462 and Theta2 = Theta3 = 2.132856
# Three "skews" based on a combination of U, V, and C(u,v) [nuskew()]
nuskewCOP(cop=GHcop, 1.538462) # zero bivariate skewness
nuskewCOP(cop=GHcop, c(2.132856, 0.6, 0.9)) # 0.1319304
nuskewCOP(cop=GHcop, c(2.132856, 0.9, 0.6)) # -0.1319304
```

So, holding τ_C constant, we see that the **GH** has a $\nu_{\text{GH}(1.538)} = 0$, but the asymmetric case $\nu_{\text{GH}(2.133,0.6,0.9)} = 0.1319304$ and $\nu_{\text{GH}(2.133,0.9,0.6)} = -0.1319304$, where the change in sign represents reflection about an equal value line. Finally, compute *L-coskew* by large simulation and the adjective “bow” representing the direction of curvature (bowing) of the principle copula density.

```
# Simulations for Theta1 would show no curvature about the diagonal.
# Simulations for Theta2 would show curvature towards the upper left.
# Simulations for Theta3 would show curvature towards the lower right.
UV <- simCOP(1000, cop=GHcop, para=c(Theta1))
UV <- simCOP(1000, cop=GHcop, para=c(Theta2, 0.6, 0.9))
UV <- simCOP(1000, cop=GHcop, para=c(Theta3, 0.9, 0.6))
# Because the Tau's are all similar, there is nothing to learn from the
# L-correlation, let us inspect the L-coskew instead:
lcm.1 <- lcomCOP(cop=GHcop, para=c(Theta1))
lcm.2 <- lcomCOP(cop=GHcop, para=c(Theta2, 0.6, 0.9))
lcm.3 <- lcomCOP(cop=GHcop, para=c(Theta3, 0.9, 0.6))
lcm.1 <- lapply(lcm.1, function(i) round(i, digits=5))
lcm.2 <- lapply(lcm.2, function(i) round(i, digits=5))
lcm.3 <- lapply(lcm.3, function(i) round(i, digits=5))

message("# T3[12]: ", lcm.1$lcomUV[3], " (symmetric) ", # L-coskews
        lcm.2$lcomUV[3], " (asym.--bow UL) ",
        lcm.3$lcomUV[3], " (asym.--bow UL)", "\n",
        "# T3[21]: ", lcm.1$lcomVU[3], " (symmetric) ",
        lcm.2$lcomVU[3], " (asym.--bow LR) ",
        lcm.3$lcomVU[3], " (asym.--bow LR)")
message("# T4[12]: ", lcm.1$lcomUV[4], " (symmetric) ", # L-cokurtoses
        lcm.2$lcomUV[4], " (asym.--bow UL) ",
        lcm.3$lcomUV[4], " (asym.--bow UL)", "\n",
        "# T4[21]: ", lcm.1$lcomVU[4], " (symmetric) ",
        lcm.2$lcomVU[4], " (asym.--bow LR) ",
        lcm.3$lcomVU[4], " (asym.--bow LR)")

# T3[12]: 0.06962 (symmetric) 0.11321 (asym.--bow UL) 0.01426 (asym.--bow UL)
# T3[21]: 0.06962 (symmetric) 0.01426 (asym.--bow LR) 0.11321 (asym.--bow LR)
```

```
# T4[12]: 0.05100 (symmetric) 0.04266 (asym.--bow UL) 0.02718 (asym.--bow UL)
# T4[21]: 0.05100 (symmetric) 0.02718 (asym.--bow LR) 0.04266 (asym.--bow LR)
```

Thus, the *L-comoments* (Asquith, 2011) using their sample values measure something fundamental about the bivariate association between the three copulas choosen. The L-coskews for the symmetrical case are equal and are

$$\tau_{3[12]}^{\mathbf{GH}(\Theta_1)} = \tau_{3[21]}^{\mathbf{GH}(\Theta_1)} = 0.06962,$$

whereas the L-coskew for the curvature to the upper left (LU) are

$$\tau_{3[12]}^{\mathbf{GH}(\Theta_2, 0.6, 0.9)} = 0.11321 \text{ and } \tau_{3[12]}^{\mathbf{GH}(\Theta_2, 0.6, 0.9)} = 0.01426,$$

whereas the L-coskew for the curvature to the lower right (LR) are

$$\tau_{3[12]}^{\mathbf{GH}(\Theta_3, 0.9, 0.6)} = 0.01426 \text{ and } \tau_{3[12]}^{\mathbf{GH}(\Theta_3, 0.6, 0.9)} = 0.11321.$$

Thus, the π_2 and π_3 parameters and L-coskew to the bivariate distribution and change the values for L-kurtoses.

Author(s)

W.H. Asquith

References

- Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.
- Brahimi, B., Chebana, F., and Necir, A., 2015, Copula representation of bivariate L-moments—A new estimation method for multiparameter two-dimensional copula models: *Statistics*, v. 49, no. 3, pp. 497–521.
- Joe, H., 2014, *Dependence modeling with copulas*: Boca Raton, CRC Press, 462 p.
- Nelsen, R.B., 2006, *An introduction to copulas*: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, *Extremes in Nature—An approach using copulas*: Springer, 289 p.
- Zhang, L., and Singh, V.P., 2007, Gumbel–Hougaard copula for trivariate rainfall frequency analysis: *Journal Hydrologic Engineering*, v. 12, Special issue—Copulas in Hydrology, pp. 409–419.

See Also

[M](#), [GLcop](#), [HRcop](#), [tEVcop](#), [rhobevCOP](#)

Examples

```
Theta    <- 2.2 # Let us see if numerical and analytical tail deps are the same.
del.lamU <- abs( taildepCOP(cop=GHcop, para=Theta)$lambdaU - (2-2^(1/Theta)) )
as.logical(del.lamU < 1E-6) # TRUE

## Not run:
```

```

# The simulations match Joe (2014, p. 72) for Gumbel-Hougaard
n <- 600; nsim <- 1000; set.seed(946) # see for reproducibility
SM <- sapply(1:nsim, function(i) { rs <- semicorCOP(cop=GHcop, para=1.35, n=n)
                                     c(rs$botleft.semicor, rs$topright.semicor) })
RhoM <- round(mean(SM[1,]), digits=3)
RhoP <- round(mean(SM[2,]), digits=3)
SE.RhoM <- round( sd(SM[1,]), digits=3)
SE.RhoP <- round( sd(SM[2,]), digits=3)
SE.RhoMP <- round( sd(SM[2,] - SM[1,]), digits=3)
# Semi-correlations (sRho) and standard errors (SEs)
message("# sRho[-]=", RhoM, " (SE[-]=", SE.RhoM, ") Joe(p.72)=0.132 (SE[-]=0.08)")
message("# sRho[+]=", RhoP, " (SE[+]=", SE.RhoP, ") Joe(p.72)=0.415 (SE[+]=0.07)")
message("# SE(sRho[-] - sRho[+])=", SE.RhoMP, " Joe(p.72) SE=0.10")
# sRho[-]=0.134 (SE[-]=0.076) Joe(p.72)=0.132 (SE[-]=0.08)
# sRho[+]=0.407 (SE[+]=0.074) Joe(p.72)=0.415 (SE[+]=0.07)
# SE(sRho[-] - sRho[+])=0.107 Joe(p.72) SE=0.10
# Joe (2014, p. 72) reports the values 0.132, 0.415, 0.08, 0.07, 0.10, respectively.
## End(Not run)

## Not run:
# Let us compare the conditional simulation method of copBasic by numerics and by the
# above analytical solution for the Gumbel-Hougaard copula to two methods implemented
# by package gumbel, a presumed Archimedean technique by package acopula, and an
# Archimedean technique by package copula. Setting seeds by each "method" below does
# not appear diagnostic because of the differences in which the simulations are made.
nsim <- 10000; kn <- "kendall" # The theoretical KENDALL TAU is (1.5-1)/1.5 = 1/3
# Simulate by conditional simulation using numerical derivative and then inversion
A <- cor(copBasic::simCOP(nsim, cop=GHcop, para=1.5, graphics=FALSE), method=kn)[1,2]
U <- runif(nsim) # GHcop.derCOPinv() defined earlier in this documentation.
V <- sapply(1:nsim, function(i) { GHcop.derCOPinv(U[i], runif(1), para=1.5) })
# Simulate by conditional simulation using exact analytical solution
B <- cor(U, y=V, method=kn); rm(U, V)
# Simulate by the "common frailty" technique
C <- cor(gumbel::rgumbel(nsim, 1.5, dim=2, method=1), method=kn)[1,2]
# Simulate by "K function" (Is the K function method, Archimedean?)
D <- cor(gumbel::rgumbel(nsim, 1.5, dim=2, method=2), method=kn)[1,2]
# Simulate by an Archimedean implementation (presumably)
E <- cor(acopula::rCopula(nsim, pars=1.5), method=kn)[1,2]
# Simulate by an Archimedean implementation
G <- cor(copula::rCopula(nsim, copula::gumbelCopula(1.5)), method=kn)[1,2]
K <- round(c(A, B, C, D, E, G), digits=5); rm(A, B, C, D, E, G, kn); tx <- ", "
message("Kendall Tau: ", K[1], tx, K[2], tx, K[3], tx, K[4], tx, K[5], tx, K[6])
# Kendall Tau: 0.32909, 0.32474, 0.33060, 0.32805, 0.32874, 0.33986 -- run 1
# Kendall Tau: 0.33357, 0.32748, 0.33563, 0.32913, 0.32732, 0.32416 -- run 2
# Kendall Tau: 0.34311, 0.33415, 0.33815, 0.33224, 0.32961, 0.33008 -- run 3
# Kendall Tau: 0.32830, 0.33573, 0.32756, 0.33401, 0.33567, 0.33182 -- nsim=50000!
# All solutions are near 1/3 and it is unknown without further study which of the
# six methods would result in the least bias and (or) sampling variability.
## End(Not run)

```

Description

Compute the measure of association known as the *Gini Gamma* γ_C (Nelsen, 2006, pp. 180–182), which is defined as

$$\gamma_C = Q(C, M) + Q(C, W),$$

where $C(u, v)$ is the copula, $M(u, v)$ is the **M** function, and $W(u, v)$ is the **W** function. The function $Q(a, b)$ (**concordCOP**) is a *concordance function* (Nelsen, 2006, p. 158). Nelsen also reports that “Gini Gamma measures a concordance relation of “distance” between $C(u, v)$ and monotone dependence, as represented by the *Fréchet–Hoeffding lower bound* and *Fréchet–Hoeffding upper bound* copulas [$M(u, v)$, **M** and $W(u, v)$, **W** respectively].”

A simpler method of computation and the default for giniCOP is to compute γ_C by

$$\gamma_C = 4 \left[\int_{\mathcal{I}} C(u, u) du + \int_{\mathcal{I}} C(u, 1 - u) du \right] - 2,$$

or in terms of the *primary diagonal* (alt. *main diagonal*, Genest *et al.*, 2010) $\delta(t)$ and *secondary diagonal* $\delta^*(t)$ (refer to **diagCOP**) by

$$\gamma_C = 4 \left[\int_{\mathcal{I}} \delta(t) dt + \int_{\mathcal{I}} \delta^*(t) dt \right] - 2.$$

The simpler method is more readily implemented because single integration is fast. Lastly, Nelsen *et al.* (2001, p. 281) show that γ_C also is computable by

$$\gamma_C = 2 Q(C, A),$$

where A is a *convex combination* (**convex2COP**, using $\alpha = 1/2$) of the copulas M and W or $A = (M + W)/2$. However, integral convergence errors seem to trigger occasionally, and the first definition by summation $Q(C, M) + Q(C, W)$ thus is used. The convex combination is demonstrated in the **Examples** section.

Usage

```
giniCOP(cop=NULL, para=NULL, by.concordance=FALSE, as.sample=FALSE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
by.concordance	Instead of using the single integrals (Nelsen, 2006, pp. 181–182) to compute γ_C , use the concordance function method implemented through concordCOP ;
as.sample	A logical controlling whether an optional R data.frame in para is used to compute the $\hat{\gamma}_C$ (refer to Note); and
...	Additional arguments to pass, which are dispatched to the copula function cop and possibly concordCOP if by.concordance=TRUE, such as delta used by that function.

Value

The value for γ_C is returned.

Note

Conceptually, the sample Gini Gamma ($\hat{\gamma}$; Genest *et al.*, 2010) is

$$\hat{\gamma} = \frac{1}{\lfloor n^2/2 \rfloor} \sum_{i=1}^n |(n+1-R_i) - S_i| - |R_i - S_i|,$$

where $\lfloor m \rfloor$ denotes the integer part of arbitrary $m > 0$, R_i and S_i are the respective ranks of X and Y and n is sample size. The sampling variance of $\hat{\gamma}$ under *assumption of independence* between X and Y is

$$\text{var}(\hat{\gamma})_{n \text{ even}} = \frac{2}{3} \frac{(n^2 + 2)}{(n-1)n^2} \text{ and}$$

$$\text{var}(\hat{\gamma})_{n \text{ odd}} = \frac{2}{3} \frac{(n^2 + 3)}{(n-1)(n^2 - 1)}.$$

Genest *et al.* (2010) present additional equations for estimation of the distribution $\hat{\gamma}$ variance for conditions of dependence based on copulas. For a copula having continuous partial derivatives, then as $n \rightarrow \infty$, the quantity $(\hat{\gamma} - \gamma_{\mathbf{C}})\sqrt{n} \sim \text{Normal}(0, \text{var}(\gamma_{\mathbf{C}}))$. Genest *et al.* (2010) show variance of $\hat{\gamma}$ for the *independence copula* ($\mathbf{C}(u, v) = \mathbf{\Pi}(u, v)$) (**P**) as $\text{var}(\gamma_{\mathbf{C}}) = 2/3$. For comparison, the *Spearman Footrule* for independence is smaller at $\text{var}(\psi_{\mathbf{C}}) = 2/5$ (refer to **footCOP Note**). In Genest *et al.* independence and two examples of dependence (p. 941), $\text{var}(\hat{\gamma}) > \text{var}(\hat{\psi})$, but those authors do not appear to remark on whether this inequality holds for all copula.

Author(s)

W.H. Asquith

References

- Genest, C., Nešlehová, J., and Ghorbal, N.B., 2010, Spearman's footrule and Gini's gamma—A review with complements: *Journal of Nonparametric Statistics*, v. 22, no. 8, pp. 937–954, [doi:10.1080/10485250903499667](https://doi.org/10.1080/10485250903499667).
- Nelsen, R.B., 2006, *An introduction to copulas*: New York, Springer, 269 p.
- Nelsen, R.B., Quesada-Molina, J.J., Rodríguez-Lallena, J.A., Úbeda-Flores, M., 2001, Distribution functions of copulas—A class of bivariate probability integral transforms: *Statistics and Probability Letters*, v. 54, no. 3, pp. 277–282, [doi:10.1016/S01677152\(01\)000608](https://doi.org/10.1016/S01677152(01)000608).

See Also

[blomCOP](#), [footCOP](#), [hoefCOP](#), [rhoCOP](#), [tauCOP](#), [wolfCOP](#), [joeskewCOP](#), [uv1moms](#)

Examples

```

giniCOP(cop=PSP)                                #                                = 0.3819757

## Not run:
giniCOP( cop=PSP, by.concordance=TRUE)           # Q(C,M) + Q(C,W) = 0.3820045
# use convex combination ---triggers integration warning but returns anyway
cxpara <- list(alpha=1/2, cop1=M, cop2=W) # parameters for convex2COP()
2*tauCOP(cop=PSP, cop2=convex2COP, para2=cxpara) # 2*Q(C,A) = 0.3819807
# where the later issued warnings on the integration
## End(Not run)

## Not run:
n <- 2000; UV <- simCOP(n=n, cop=N4212cop, para=9.3, graphics=FALSE)
giniCOP(para=UV, as.sample=TRUE)                 # 0.9475900 (sample version)
giniCOP(cop=N4212cop, para=9.3)                  # 0.9479528 (copula integration)
giniCOP(cop=N4212cop, para=9.3, by.concordance=TRUE) # 0.9480267 (concordance func.)
# where the later issued warnings on the integration
## End(Not run)

## Not run:
# The canonical example of theoretical and sample estimators of bivariate
# association for the package: Blomqvist Beta, Spearman Footrule, Gini Gamma,
# Hoeffding Phi, Kendall Tau, Spearman Rho, and Schweizer-Wolff Sigma
# and comparison to L-correlation via lmomco::lcomoms2().
n <- 9000; set.seed(56)
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, para1=1.45, para2=21.9,
             alpha=0.41, beta=0.08)
D <- simCOP(n=n, cop=composite2COP, para=para, cex=0.5, col=rgb(0,0,0.2), pch=16)
blomCOP(cop=composite2COP, para=para)            # 0.4037908 (theoretical)
blomCOP(para=D, as.sample=TRUE)                  # 0.4008889 (sample)
footCOP(cop=composite2COP, para=para)            # 0.3721555 (theoretical)
footCOP(para=D, as.sample=TRUE)                  # 0.3703623 (sample)
giniCOP(cop=composite2COP, para=para)            # 0.4334687 (theoretical)
giniCOP(para=D, as.sample=TRUE)                  # 0.4311698 (sample)
tauCOP(cop=composite2COP, para=para)             # 0.3806909 (theoretical)
tauCOP(para=D, as.sample=TRUE)                   # 0.3788139 (sample)
rhoCOP(cop=composite2COP, para=para)             # 0.5257662 (theoretical)
rhoCOP(para=D, as.sample=TRUE)                   # 0.5242380 (sample)
lmomco::lcomoms2(D)$T2                          # 1
                                                  # 0.5242388 (sample matrix)
                                                  # 0.5245154 1
hoefCOP(cop=composite2COP, para=para)            # 0.5082776 (theoretical)
subsample <- D[sample(1:n, n/5),] # subsampling for speed
hoefCOP(para=subsample, as.sample=TRUE)          # 0.5033842 (re-sample)
#hoefCOP(para=D, as.sample=TRUE) # major CPU hog, n too big
# because the Ds are already "probabilities" just resample as shown above
wolfCOP(cop=composite2COP, para=para)            # 0.5257662 (theoretical)
wolfCOP(para=D, as.sample=TRUE)                  # 0.524239 (by fast grid)
wolfCOP(para=subsample, as.sample=TRUE)          # 0.5338009 (re-sample)
## End(Not run)

## Not run:
set.seed(1); nsim <- 1000

```

```

varGIunderIndpendence <- function(n) {
  if(3 %% 2 == 3 / 2) { # even
    (2/3)*( (n^2 + 2) ) / ( (n-1)* n^2 ) # Genest et al. (2010)
  } else {
    (2/3)*( (n^2 + 3) ) / ( (n-1)*(n^2-1) ) # Genest et al. (2010)
  }
}
ns <- c(10, 15, 20, 25, 30, 50, 75, 100); ZZ <- NULL
plot(min(ns):max(ns), varGIunderIndpendence(10:max(ns)), type="l", ylim=c(0,0.1),
     xlab="Sample size", ylab="Variance of Sample Estimator", col="darkgreen")
mtext("Sample Gini Gamma Under Independence", col="darkgreen", line=1)
mtext("Sample Schweizer-Wolff Sigma Under Independence", col="salmon4")
for(n in ns) {
  sGI <- sSW <- vector(length=nsim)
  for(i in seq_len(nsim)) {
    uv <- simCOP(n=n, cop=P, para=2, graphics=FALSE)
    sGI[i] <- giniCOP(para=uv, as.sample=TRUE)
    sSW[i] <- wolfCOP(para=uv, as.sample=TRUE)
  }
  varGI <- varGIunderIndpendence(n)
  zz <- round(c(n, mean(sGI), var(sGI), varGI, mean(sSW), var(sSW)), digits=6)
  names(zz) <- c("n", "mean_sim", "var_sim", "var_Genest", "mean_Wolf", "var_Wolf")
  print(zz); ZZ <- rbind(ZZ, data.frame(t(zz)))
  points(n, zz[3], cex=2, pch=21, col="darkgreen", bg="lightgreen")
  points(n, zz[6], cex=2, pch=22, col="salmon4", bg="salmon1" )
} # results show proper implementation and Genest et al. (2010, sec. 3) with the
# Schweizer-Wolff comparison unique to this copBasic Example.
## End(Not run)

```

GLcop

The Galambos Extreme Value Copula (with Gamma Power Mixture [Joe/BB4] and Lower Extreme Value Limit)

Description

The *Galambos copula* (Joe, 2014, p. 174) is

$$\mathbf{C}_{\Theta}(u, v) = \mathbf{GL}(u, v) = uv \exp \left[\left(x^{-\Theta} + y^{-\Theta} \right)^{-1/\Theta} \right],$$

where $\Theta \in [0, \infty)$, $x = -\log u$, and $y = -\log v$. As $\Theta \rightarrow 0^+$, the copula limits to *independence* (**II**; **P**) and as $\Theta \rightarrow \infty$, the copula limits to perfect association (**M**; **M**). The Galambos is a *bivariate extreme value copula* (BEV), and parameter estimation for Θ requires numerical methods.

There are two other genetically related forms. Joe (2014, p. 197) describes an extension of the Galambos copula as a *Galambos gamma power mixture* (GLPM), which is Joe's *BB4 copula*, with the following form

$$\mathbf{C}_{\Theta, \delta}(u, v) = \mathbf{GLPM}(u, v) = \left(x + y - 1 - [(x-1)^{-\delta} + (y-1)^{-\delta}]^{-1/\delta} \right)^{-1/\Theta},$$

where $x = u^{-\Theta}$, $y = v^{-\Theta}$, and $\Theta \geq 0, \delta \geq 0$. (Joe shows $\delta > 0$, but zero itself seems to work without numerical problems in practical application.) As $\delta \rightarrow 0^+$, the “MTCJ family” (Mardia–Takahasi–Cook–Johnson) results (implemented internally with Θ as the incoming parameter). As $\Theta \rightarrow 0^+$, the Galambos above results with δ as the incoming parameter.

This Galambos extension in turn has a *lower extreme value limit form* that leads to a *min-stable bivariate exponential* having *Pickand dependence function* of

$$A(x, y; \Theta, \delta) = x + y - \left[x^{-\Theta} + y^{-\Theta} - (x^{\Theta\delta} + y^{\Theta\delta})^{-1/\delta} \right]^{-1/\Theta},$$

where this third copula is

$$\mathbf{C}_{\Theta, \delta}^{LEV}(u, v) = \mathbf{GLEV}(u, v) = \exp[-A(-\log u, -\log v; \Theta, \delta)],$$

for $\Theta \geq 0, \delta \geq 0$ and is known as the *two-parameter Galambos*. (Joe shows $\delta > 0$, but $\delta = 0$ itself seems to work without numerical problems in practical application.)

Usage

```
GLcop( u, v, para=NULL, ...)
GLEVcop( u, v, para=NULL, ...)
GLPMcop( u, v, para=NULL, ...) # inserts third parameter automatically
J0copBB4(u, v, para=NULL, ...) # inserts third parameter automatically
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	To trigger $\mathbf{GL}(u, v)$, a vector (single element) of Θ , to trigger $\mathbf{GLEV}(u, v)$, a two element vector of Θ and δ and alias is <code>GLEVcop</code> , and to trigger $\mathbf{GLPM}(u, v)$, a three element vector of Θ , δ , and any number (the presence of the third entry alone is the triggering mechanism) though aliases <code>GLPM</code> or <code>J0copBB4</code> will insert the third parameter automatically for convenience; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Note

Joe (2014, p. 198) shows $\mathbf{GLEV}(u, v; \Theta, \delta)$ as a two-parameter Galambos, but its use within the text seemingly is not otherwise obvious. However, testing of the implementation here seems to show that this copula is really not broader in form than $\mathbf{GL}(u, v; \alpha)$. The α can always(?) be chosen to mimic the $\{\Theta, \delta\}$. This assertion can be tested from a semi-independent direction. First, define an alternative style of one-parameter Galambos:

```

GL1cop <- function(u,v, para=NULL, ...) {
  GL1pA <- function(x,y,t) { # Pickend dependence func form 1p Galambos
    x + y - (x^-t + y^-t)^(-1/t)
  }
  if(length(u) == 1) { u <- rep(u, length(v)) } else
  if(length(v) == 1) { v <- rep(v, length(u)) }
  exp(-GL1pA(-log(u), -log(v), para[1]))
}

```

Second, redefine the two-parameter Galambos:

```

GL2cop <- function(u,v, para=NULL, ...) {
  GL2pA <- function(x,y,t,d) { # Pickend dependence func form 2p Galambos
    x + y - (x^-t + y^-t - (x^(t*d) + y^(t*d))^(-1/d))^(-1/t)
  }
  if(length(u) == 1) { u <- rep(u, length(v)) } else
  if(length(v) == 1) { v <- rep(v, length(u)) }
  exp(-GL2pA(-log(u), -log(v), para[1], para[2]))
}

```

Next, we can combine the Pickend dependence functions into an objective function. This objective function will permit the computation of the α given a pair $\{\Theta, \delta\}$.

```

objfunc <- function(a,t=NA,d=NA, x=0.7, y=0.7) {
  lhs <- (x^-t + y^-t - (x^(t*d) + y^(t*d))^(-1/d))^(-1/t)
  rhs <- (x^-a + y^-a)^(-1/a); return(rhs - lhs) # to be uniroot'ed
}

```

A demonstration can now be made:

```

t <- 0.6; d <- 4; lohi <- c(0,100)
set.seed(3); UV <- simCOP(3000, cop=GL2cop, para=c(t,d), pch=16,col=3,cex=0.5)
a <- uniroot(objfunc, interval=lohi, t=t, d=d)$root
set.seed(3); UV <- simCOP(3000, cop=GL1cop, para=a, lwd=0.5, ploton=FALSE)

```

The graphic so produced shows almost perfect overlap in the simulated values. To date, the author has not really found that the two parameters can be chosen such that the one-parameter version can not attain. Extensive numerical experiments using simulated parameter combinations through the use of various copula metrics (tail dependencies, L-comoments, etc) have not found material differences. **Has the author of this package missed something?**

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[M](#), [P](#), [GHcop](#), [HRcop](#), [tEvcop](#)

Examples

```
# Theta = pi for GLcop and recovery through Blomqvist Beta      (Joe, 2014, p. 175)
log(2)/(log(log(2)/log(1+blomCOP(cop=GLcop, para=pi))))

# Theta = 2 and delta = 3 for the GLPM form and Blomqvist Beta (Joe, 2014, p. 197)
t <- 2; Btheo <- blomCOP(GLPMcop, para=c(t,3))
Bform <- (2^(t+1) - 1 - taildepCOP(GLPMcop, para=c(t,3))$lambdaU*(2^t -1))^(1/t)
print(c(Btheo, 4*Bform-1)) # [1] 0.8611903 0.8611900

## Not run:
# See the Note section but check Blomqvist Beta here:
blomCOP(cop=GLcop, para=c(6.043619)) # 0.8552863 (2p version)
blomCOP(cop=GLcop, para=c(5.6, 0.3)) # 0.8552863 (1p version)
## End(Not run)
```

glueCOP

Gluing Two Copulas

Description

The *gluing copula* technique (Erdely, 2017, p. 71), given two bivariate copulas $\mathbf{C}_A$ and $\mathbf{C}_B$ and a fixed value $0 \leq \gamma \leq 1$ is

$$\mathbf{C}_\gamma(u, v) = \gamma \cdot \mathbf{C}_A(u/\gamma, v)$$

for $0 \leq u \leq \gamma$ and

$$\mathbf{C}_\gamma(u, v) = (1 - \gamma) \cdot \mathbf{C}_B((u - \gamma) / (1 - \gamma), v)$$

for $\gamma \leq u \leq 1$ and γ represents the *gluing point* in u (horizontal axis). The logic is simply the rescaling of $\mathbf{C}_A$ to $[0, \gamma] \times [0, 1]$ and $\mathbf{C}_B$ to $[\gamma, 1] \times [0, 1]$. Copula gluing is potentially useful in circumstances for which regression is non-monotone.

Usage

```
glueCOP(u, v, para=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A special parameter list (see Note) with a mandatory element of glue parameter γ ; and
<code>...</code>	Additional arguments to pass to the copulas.

Value

Value(s) for the copula are returned.

Note

The following descriptions list in detail the structure and content of the `para` argument:

`glue` — The γ gluing parameter;
`cop1` — Function of the first copula **A**;
`cop2` — Function of the second copula **B**;
`para1` — Vector of parameters $\Theta_{\mathbf{A}}$ for **A**; and
`para2` — Vector of parameters $\Theta_{\mathbf{B}}$ for **B**.

Author(s)

W.H. Asquith

References

Erdely, A., 2017, Copula-based piecewise regression (chap. 5) *in* Copulas and dependence models with applications—Contributions in honor of Roger B. Nelsen, *eds.* Flores, U.M., Amo Artero, E., Durante, F., Sánchez, J.F.: Springer, Cham, Switzerland, ISBN 978–3–319–64220–9, doi:10.1007/9783319642215.

See Also

[COP](#), [breveCOP](#), [composite1COP](#), [composite2COP](#), [composite3COP](#), [convexCOP](#)

Examples

```
## Not run:
para <- list(cop1=PLACKETTcop, para1=.2, cop2=GLcop, para2=1.2, glue=0.6)
densityCOPplot(cop=glueCOP, para=para) #
## End(Not run)

## Not run:
# Concerning Nelsen (2006, exam. 3.3, pp. 59-61)
# Concerning Erdely (2017, exam. 5.1, p. 71)
# Concerning Erdely (2017, exam. 5.2, p. 75)
# Nelsen's example is a triangle with vertex at [G, 1].
# Erdely's example permits the construction using glueCOP from M and W.
"coptri" <- function(u,v, para=NA, ...) {
  p <- para[1]; r <- 1 - (1-p)*v
  if(length(u) > 1 | length(v) > 1) stop("only scalars for this function")
  if(0 <= u & u <= p*v & p*v <= p) { return(u)
  } else if( 0 <= p*v & p*v < u & u < r) { return(p*v)
  } else if( p <= r & r <= u & u <= 1 ) { return(u+v-1)
  } else { stop("should not be here in logic") }
}
"UsersCOP" <- function(u,v, ...) { asCOP(u,v, f=coptri, ...) }
```

```

# Demonstrate Nelsen's triangular copula (black dots )
UV <- simCOP(cop=UsersCop, para=0.35, cex=0.5, pch=16)
# Add Erdley's gluing of M() and W() copula (red circles)
para <- list(cop1=M, cop2=W, para1=NA, para2=NA, glue=0.35)
UV <- simCOP(cop=glueCOP, para=para, col=2, ploton=FALSE)
# We see in the plot that the triangular copulas are the same.

# For G = 0.5, Erdley shows Spearman Rho = 2*G-1 = 0, but
# Schweizer-Wolff = G^2 + (G-1)^2 = 0.5, let us check these:
para <- list(cop1=M, cop2=W, para1=NA, para2=NA, glue=0.5)
rhoCOP( cop=glueCOP, para=para) # -2.181726e-17
wolfCOP(cop=glueCOP, para=para) # 0.4999953
# So, rhoCOP() indicates independence, but wolfCOP() indicates
# dependence at the minimum value possible for a triangular copula.
## End(Not run)

## Not run:
x <- seq(0,1, by=0.05); para <- list(cop1=M, cop2=W, para1=NA, para2=NA)
LCM <- lapply(x, function(i) { para$glue <- i;
  lcomCOP(cop=glueCOP, para=para, stop.on.error=FALSE) })
LCM12 <- lapply(LCM, function(i) i$lcomUV)
LCM21 <- lapply(LCM, function(i) i$lcomVU)
LCM12 <- t( sapply(LCM12, function(i) i) )
LCM21 <- t( sapply(LCM21, function(i) i) )
plot( x, LCM12[,3], ylim=c(-0.8, +0.8), type="l", lty=2,
  xlab="glueCOP() gluing parameter value",
  ylab="L-coskew or L-cokurtosis")
lines(x, LCM21[,3], col="black")
lines(x, LCM12[,4])
lines(x, LCM21[,4], col="darkorchid3")
rho <- LCM12[,2]
plot(rho, LCM12[,3], ylim=c(-0.8, +0.8), type="l", lty=2,
  xlab="Spearman Rho", ylab="L-coskew or L-cokurtosis")
lines(rho, LCM21[,3], col="black")
lines(rho, LCM12[,4], lty=2)
lines(rho, LCM21[,4], col="darkorchid3") #
## End(Not run)

```

gridCOP

Compute a Copula on a Grid

Description

Compute a grid of copula values. This function has the primary intention of supporting 3D renderings or 2D images of the *copulatic surface*. Users should be aware of the convention of the placement of the plotting origin and the various plotting mechanisms available to them in R. By convention copulatic surfaces start in lower left corner for $u = v = 0$, but matrix conventions (or at least how some functions plot matrices) start with the origin in the upper left.

Usage

```
gridCOP(cop=NULL, para=NULL, delta=0.05, transpose=TRUE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
delta	The $\Delta u = \Delta v$ of the grid edges;
transpose	A logical to transpose the returned grid. This is needed if functions such as <code>image()</code> in R are to be used for visualization (see last example in Examples with composite2COP); and
...	Additional arguments to pass.

Value

The values for $C(u, v)$ are returned as a grid as an R matrix.

Author(s)

W.H. Asquith

See Also

[EMPIRcopdf](#)

Examples

```
## Not run:
the.grid <- gridCOP(cop=PSP)
the.grid[1,1] <- 0 # replace the NaN
image(the.grid) # ramps to the upper right
## End(Not run)

## Not run:
# See this composite copula also used in densityCOPplot() documentation.
para <- list(alpha=0.15, beta=0.90, kappa=0.06, gamma=0.96,
             cop1=GHcop, cop2=PLACKETTcop, para1=5.5, para2=0.07)
GR <- gridCOP(cop=composite2COP, para=para, delta=0.005)
image(GR, col=terrain.colors(20)) # asymmetric, high curvature in top half
## End(Not run)
```

HAIRPINcop	<i>The Hairpin Copula</i>
------------	---------------------------

Description

The *Hairpin copula* (Durante *et al.*, 2014) is

$$\mathbf{C}_{\Theta}(u, v) = \mathbf{HAIRPIN}(u, v) = \min[u, v, \text{mean}(f(u, v))],$$

as vectorized for the pairs (u_i, v_i) for $i \in 1, 2, \dots$, and $f(x) = \alpha x^{\phi}$, $1 \leq \alpha \leq \infty$, and $1 \leq \phi \leq 2$. The copula, as $\alpha \rightarrow \infty$ or $\phi \rightarrow 1$ limits, to the *comonotonicity coupla* ($\mathbf{M}(u, v)$; [M](#)).

Usage

HAIRPINcop(u, v, para=c(1, 2, 1), ...)

Arguments

- | | |
|------|--|
| u | Nonexceedance probability u in the X direction; |
| v | Nonexceedance probability v in the Y direction; |
| para | A vector (three element) of two parameters and the reflection: α and ϕ of the coupla then the reflection number $n \in 1, 2, 3, 4$ following the reflection pattern of COP ; and |
| ... | Additional arguments to pass. |

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Durante, F., Fernández-Sánchez, J., and Trutschnig, W., 2014, Multivariate copulas with hairpin support: Journal of Multivariate Analysis, v. 130, pp. 323–334, doi:[10.1016/j.jmva.2014.06.009](https://doi.org/10.1016/j.jmva.2014.06.009).

See Also

[M](#)

Examples

```
## Not run:
# Produce a Celtic Knot look through simulation -----
nsim <- 1.5E4
plot(c(0, 1), c(0, 1), type="n", bty="n",
     xaxt="n", yaxt="n", xaxs="r", yaxs="r", xlab="", ylab="")
sapply(1:4, function(n) {
  UV <- simCOP(nsim, cop=HAIRPINcop, para=c(1.12, 1.8, n),
               seed=1, ploton=FALSE, pch=16, cex=0.6, col=n); return(n) })
sapply(1:4, function(n) {
  UV <- simCOP(nsim, cop=HAIRPINcop, para=c(1.12, 1.3, n),
               seed=1, ploton=FALSE, pch=16, cex=0.6, col=n); return(n) }) #
## End(Not run)
```

hoefCOP

The Hoeffding Phi of a Copula or Lp Distances (Independence, Radial Asymmetry, or Reflection Symmetry Forms)

Description

Compute the measure of association known as the *Hoeffding Phi* $\Phi_{\mathbf{C}}$ of a copula from *independence* ($uv = \mathbf{\Pi}; \mathbf{P}$) according to Cherunbini *et al.* (2004, p. 164) by

$$\Phi_{\mathbf{C}} = 3 \sqrt{10 \int \int_{\mathcal{I}^2} (\mathbf{C}(u, v) - uv)^2 \, dudv},$$

and Nelsen (2006, p. 210) shows this as (and absolute value notation by Nelsen helps in generalization)

$$\Phi_{\mathbf{C}} = \left(90 \int \int_{\mathcal{I}^2} |\mathbf{C}(u, v) - uv|^2 \, dudv \right)^{1/2},$$

for which $\Phi_{\mathbf{C}}^2$ (the square of the quantity) is known as the *dependence index*. Gaißer *et al.* (2010, eq. 1) have $\Phi_{\mathbf{C}}^2$ as the *Hoeffding Phi-Square*, and their definition, when square-rooted, matches Nelsen's listing.

A generalization (Nelsen, 2006) to L_p distances from independence ($uv = \mathbf{\Pi}; \mathbf{P}$) through the LpCOP function is

$$L_p \equiv \Phi_{\mathbf{C}}(p) = \left(k(p) \int \int_{\mathcal{I}^2} |\mathbf{C}(u, v) - uv|^p \, dudv \right)^{1/p},$$

for a $p : 1 \leq p \leq \infty$ and where $k(p)$ is a normalization constant such that $\Phi_{\mathbf{C}}(p) = 1$ when the copula $\mathbf{C}$ is $\mathbf{M}$ (see [M](#)) or $\mathbf{W}$ (see [W](#)). The $k(p)$ (bivariate definition only) for other powers is given (Nelsen, 2006, exer. 5.44, p. 213) in terms of the *complete gamma function* $\Gamma(t)$ by

$$k(p) = \frac{\Gamma(2p + 3)}{2[\Gamma(p + 1)]^2},$$

which is implemented by the `hoefCOP` function. It is important to realize that the L_p distances are all *symmetric nonparametric measures of dependence* (Nelsen, 2006, p. 210). These are symmetric because distance from independence is used as evident by “ uv ” in the above definitions.

Reflection/Radial and Permutation Asymmetry—Asymmetric forms similar to the above distances exist. Joe (2014, p. 65) shows two measures of bivariate *reflection asymmetry* or *radial asymmetry* (term favored in **copBasic**) as the distance between $\mathbf{C}(u, v)$ and the survival copula $\hat{\mathbf{C}}(u, v)$ (**surCOP**) measured by

$$L_{\infty}^{\text{radsym}} = \sup_{0 \leq u, v \leq 1} |\mathbf{C}(u, v) - \hat{\mathbf{C}}(u, v)|,$$

or its L_p^{radsym} counterpart

$$L_p^{\text{radsym}} = \left[\int \int_{T^2} |\mathbf{C}(u, v) - \hat{\mathbf{C}}(u, v)|^p \, du \, dv \right]^{1/p} \text{ with } p \geq 1,$$

where $\hat{\mathbf{C}}(u, v) = u + v - 1 + \mathbf{C}(1 - u, 1 - v)$ and again $p : 1 \leq p \leq \infty$. Joe (2014) does not appear to discuss and normalization constants for these two radial asymmetry distances.

Joe (2014, p. 66) offers analogous measures of bivariate *permutation asymmetry* (**isCOP.permsym**) ($\mathbf{C}(u, v) \neq \mathbf{C}(v, u)$) defined as

$$L_{\infty}^{\text{permsym}} = \sup_{0 \leq u, v \leq 1} |\mathbf{C}(u, v) - \hat{\mathbf{C}}(v, u)|,$$

or its L_p^{permsym} counterpart

$$L_p^{\text{permsym}} = \left[\int \int_{T^2} |\mathbf{C}(u, v) - \hat{\mathbf{C}}(v, u)|^p \, du \, dv \right]^{1/p} \text{ with } p \geq 1,$$

where $p : 1 \leq p \leq \infty$. Again, Joe (2014) does not appear to discuss and normalization constants for these two permutation symmetry distances. Joe (2014, p. 65) states that the “simplest one-parameter bivariate copula families [and] most of the commonly used two-parameter bivariate copula families are permutation symmetric.” The $L_{\infty}^{\text{permsym}}$ (or rather a similar form) is implemented by **LzCOPpermsym** and demonstration made in that documentation.

The asymmetrical L_{∞} and L_p measures identified by Joe (2014, p. 66) are nonnegative with an upper bounds that depends on p . The bound dependence on p is caused by the lack of normalization constant $k(p)$. In an earlier paragraph, Joe (2014) indicates an upper bounds of 1/3 for both (likely?) concerning $L_{\infty}^{\text{radsym}}$ and $L_{\infty}^{\text{permsym}}$. Discussion of this 1/3 or rather the integer 3 is made within **LzCOPpermsym**.

The numerical integrations for L_p^{radsym} and L_p^{permsym} can readily return zeros. Often inspection of the formula for the $\mathbf{C}(u, v)$ itself would be sufficient to judge whether symmetry exists and hence the distances are uniquely zero.

Joe (2014, p. 66) completes the asymmetry discussion with three definitions of skewness of combinations of random variables U and V : Two definitions are in **uVlmoms** (for $U + V - 1$ and $U - V$) and two are for $V - U$ (**nuskewCOP**) and $U + V - 1$ (**nustarCOP**).

Usage

```

hoefCOP(      cop=NULL, para=NULL, p=2, as.sample=FALSE,
              sample.as.prob=TRUE,
              brute=FALSE, delta=0.002, ...)

LpCOP(      cop=NULL, para=NULL, p=2, brute=FALSE, delta=0.002, ...)
LpCOPradsym( cop=NULL, para=NULL, p=2, brute=FALSE, delta=0.002, ...)
LpCOPpermsym(cop=NULL, para=NULL, p=2, brute=FALSE, delta=0.002, ...)

```

Arguments

<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>p</code>	The value for p as described above with a default to 2 to match the discussion of Nelsen (2006) and the <i>Hoeffding Phi</i> of Cherubini <i>et al.</i> (2004). Do not confuse p with d described in Note ;
<code>as.sample</code>	A logical controlling whether an optional R data.frame in <code>para</code> is used to compute the $\hat{\Phi}_C$ (see Note). If set to -1, then the message concerning CPU effort will be suppressed;
<code>sample.as.prob</code>	When <code>as.sample</code> triggered, what are the units incoming in <code>para</code> ? If they are probabilities, the default is applicable. If they are not, then the columns are re-ranked and divided simply by $1/n$ —more sophisticated <i>empirical copula</i> probabilities are not used (EMPIRCop);
<code>brute</code>	Should brute force be used instead of two nested <code>integrate()</code> functions in R to perform the double integration;
<code>delta</code>	The du and dv for the brute force (<code>brute=TRUE</code>) integration; and
<code>...</code>	Additional arguments to pass.

Value

The value for $\Phi_C(p)$ is returned.

Note

Concerning the distance from independence, when $p = 1$, then the *Spearman Rho* ([rhoCOP](#)) of a copula is computed where it is seen in that documentation that the $k_p(1) = 12$. The respective values of $k(p)$ for select integers p are

$$p \mapsto [1, 2, 3, 4, 5] \equiv k(p) \mapsto \{12, 90, 560, 3150, 16600\},$$

and these values are hardwired into `hoefCOP` and `LpCOP`. The integers for k_p ensures that the equality in the second line of the examples is TRUE, but the p can be a noninteger as well. Nelsen (2006, p. 211) reports that when $p = \infty$ that L_∞ is

$$L_\infty \equiv \Phi_C(\infty) = \Lambda_C = 4 \sup_{u,v \in \mathcal{I}} |\mathbf{C}(u, v) - uv|.$$

A sample $\hat{\Phi}_{\mathbf{C}}$ (square root of the *Hoeffding Phi-Square*) based on nonparametric estimation generalized for d dimensions ($d = 2$ for bivariate) is presented by Gaißer *et al.* (2010, eq. 10) for estimated probabilities $\hat{U}_{ij}$ for the i th dimension and j th row (observation) for sample of size n . Those authors suggest that the $\hat{U}_{ij}$ be estimated from the empirical copula:

$$\hat{\Phi}_{\mathbf{C}} = \sqrt{h(d)[A + B]},$$

where

$$A = \left(\frac{1}{n}\right)^2 \sum_{j=1}^n \sum_{k=1}^n \prod_{i=1}^d [1 - \max(\hat{U}_{ij}, \hat{U}_{ik})],$$

$$B = \left(\frac{1}{3}\right)^d - \left(\frac{2}{n}\right) \left(\frac{1}{2}\right)^d \sum_{j=1}^n \prod_{i=1}^d [1 - \hat{U}_{ij}^2].$$

The normalization constant is a function of dimension and is

$$h(d)^{-1} = \frac{2}{(d+1)(d+2)} - \left(\frac{1}{2}\right)^d \frac{d!}{\prod_{i=0}^d (i + (1/2))} + \left(\frac{1}{3}\right)^d.$$

```
set.seed(1); UV <- simCOP(n=1000, cop=PSP)
hoefCOP(cop=PSP) # 0.4547656 (theo.)
hoefCOP(para=UV, as.sample=TRUE) # 0.4892757
set.seed(1); UV <- simCOP(n=1000, cop=PSP, snv=TRUE) # std normal variates
hoefCOP(para=UV, as.sample=TRUE, sample.as.prob=FALSE) # 0.4270324
```

Author(s)

W.H. Asquith

References

- Cherubini, U., Luciano, E., and Vecchiato, W., 2004, Copula methods in finance: Hoboken, NJ, Wiley, 293 p.
- Gaißer, S., Ruppert, M., and Schmid, F., 2010, A multivariate version of Hoeffding's Phi-Square: Journal of Multivariate Analysis, v. 101, no. 10, pp. 2571–2586, doi:10.1016/j.jmva.2010.07.006.
- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

blomCOP, footCOP, giniCOP, rhoCOP, tauCOP, wolfCOP, joeskewCOP, uvlmoms, LzCOPpermsym

Examples

```
## Not run:
# Example (ii) Gaißer et al. (2010, p. 2574)
Theta <- 0.66 # Phi^2 = Theta^2 ---> Phi == Theta as shown
hoefCOP(cop=convex2COP, para=c(alpha=Theta, cop1=M, cop2=P)) # 0.6599886

rhoCOP(cop=PSP) == hoefCOP(cop=PSP, p=1) # TRUE
LpCOP(cop=PLACKETTcop, para=1.6, p=2.6) # 0.1445137 (Fractional p)
## End(Not run)

## Not run:
set.seed(938) # Phi(1.6; Plackett) = 0.1184489; L_1 = 0.1168737
UV <- simCOP(cop=PLACKETTcop, para=1.6, n=2000, ploton=FALSE, points=FALSE)
hoefCOP(cop=PLACKETTcop, para=1.6, p=200) # Large p near internal limits
L_1 <- 4*max(abs(PLACKETTcop(UV$U, UV$V, para=1.6) - UV$U*UV$V)) # p is infy
# and finite n and arguably a sample-like statistic here, now on intuition try
# a more sample-like means
U <- runif(10000); V <- runif(10000)
L_2 <- 4*max(abs(EMPIRCOP(U, V, para=UV) - U*V)) # 0.1410254 (not close enough)
## End(Not run)

## Not run:
para <- list(alpha=0.15, beta=0.90, kappa=0.06, gamma=0.96,
             cop1=GHcop, cop2=PLACKETTcop, para1=5.5, para2=0.07)
LpCOPradsym( cop=composite2COP, para=para) # 0.02071164
LpCOPpermsym(cop=composite2COP, para=para) # 0.01540297
## End(Not run)

## Not run:
"M0cop.formula" <- function(u,v, para=para, ...) {
  alpha <- para[1]; beta <- para[2]; return(min(v*u^(1-alpha), u*v^(1-beta)))
}
"M0cop" <- function(u,v, ...) { asCOP(u,v, f=M0cop.formula, ...) }
LpCOPradsym( cop=M0cop, para=c(0.8, 0.5)) # 0.0261843
LpCOPpermsym(cop=M0cop, para=c(0.8, 0.5)) # 0.0243912
## End(Not run)
```

Description

The *Hüsler–Reiss copula* (Joe, 2014, p. 176) is

$$C_{\Theta}(u, v) = \mathbf{HR}(u, v) = \exp[-x\Phi(X) - y\Phi(Y)],$$

where $\Theta \geq 0$, $x = -\log(u)$, $y = -\log(v)$, $\Phi(\cdot)$ is the cumulative distribution function of the standard normal distribution, X and Y are defined as:

$$X = \frac{1}{\Theta} + \frac{\Theta}{2} \log\left(\frac{x}{y}\right) \text{ and } Y = \frac{1}{\Theta} + \frac{\Theta}{2} \log\left(\frac{y}{x}\right).$$

As $\Theta \rightarrow 0^+$, the copula limits to *independence* (**II**; **P**). The **HR** copula is a *bivariate extreme value copula* (*BEV*), and the parameter Θ requires numerical methods. Because there is “hardly any practical difference among [these] bivariate exchangeable parametric families” distinguishing between these (**GHcop**, **GLcop**, and **HRcop**) “requires extremely large sample sizes” (Hofert *et al.*, 2016, p. 117). This observation is strongly supported by the *L-comoment ratio diagrams* explored in **LCOMDIA_ManyCops**.

Usage

```
HRcop(u, v, para=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Hofert, M., Kojadinovic, I., Mächler, M., and Yan, J., 2018, Elements of copula modeling with R: Dordrecht, Netherlands, Springer.

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[P](#), [GHcop](#), [GLcop](#), [tEvcop](#)

Examples

```
# Parameter Theta = pi recovery through the Blomqvist Beta (Joe, 2014, p. 176)
qnorm(1 - log( 1 + blomCOP(cop=HRcop, para=pi) ) / ( 2 * log(2) ) )^(-1)
```

isCOP.LTD

*Is a Copula Left-Tail Decreasing***Description**

Numerically set a logical whether a copula is *left-tail decreasing* (LTD) as described by Nelsen (2006, pp. 192–193) and Salvadori *et al.* (2007, p. 222). A copula $\mathbf{C}(u, v)$ is left-tail decreasing for $\text{LTD}(V|U)$ if and only if for any $v \in [0, 1]$ that the following holds

$$\frac{\delta \mathbf{C}(u, v)}{\delta u} \leq \frac{\mathbf{C}(u, v)}{u}$$

for almost all $u \in [0, 1]$. Similarly, a copula $\mathbf{C}(u, v)$ is left-tail decreasing for $\text{LTD}(U|V)$ if and only if for any $u \in [0, 1]$ that the following holds

$$\frac{\delta \mathbf{C}(u, v)}{\delta v} \leq \frac{\mathbf{C}(u, v)}{v}$$

for almost all $v \in [0, 1]$ where the later definition is controlled by the `wrtV=TRUE` argument.

The LTD concept is associated with the concept of *tail monotonicity* (Nelsen, 2006, p. 191). Specifically, but reference to Nelsen (2006) definitions and geometric interpretations is recommended, $\text{LTD}(V|U)$ (or $\text{LTD}(V|U)$) means that the probability $P[Y \leq y | X \leq x]$ (or $P[X \leq x | Y \leq y]$) is a nonincreasing function of x (or y) for all y (or x).

A positive LTD of either $\text{LTD}(V|U)$ or $\text{LTD}(U|V)$ implies positively quadrant dependency (PQD, [isCOP.PQD](#)) but the condition of PQD does not imply LTD. Finally, the accuracy of the numerical assessment of the returned logical by `isCOP.LTD` is dependent on the the “smallness” of the delta argument passed into the function.

Usage

```
isCOP.LTD(cop=NULL, para=NULL, wrtV=FALSE, delta=0.005, ...)
```

Arguments

<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters, if needed, to pass to the copula;
<code>wrtV</code>	A logical to toggle between with respect to v or u (default);
<code>delta</code>	The increment of $\{u, v\} \mapsto [0 + \Delta\delta, 1 - \Delta\delta, \Delta\delta]$ set by <code>wrtV</code> ; and
<code>...</code>	Additional arguments to pass to the copula or derivative of a copula function.

Value

A logical TRUE or FALSE is returned.

Author(s)

W.H. Asquith

References

- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, Extremes in nature—An approach using copulas: Dordrecht, Netherlands, Springer, Water Science and Technology Library 56, 292 p.

See Also

[isCOP.RTI](#), [isCOP.PQD](#)

Examples

```
## Not run:
isCOP.LTD(cop=P, delta=0.01) # independence should be FALSE
# Positive association
isCOP.LTD(cop=PSP) # TRUE
# Negative association Plackett
isCOP.LTD(cop=PLACKETTcop, para=0.15) # FALSE
# Positive association Plackett
isCOP.LTD(cop=PLACKETTcop, para=15) # TRUE
# Negative association Plackett
isCOP.LTD(cop=PLACKETTcop, wrtv=TRUE, para=0.15) # FALSE
# Positive association Plackett
isCOP.LTD(cop=PLACKETTcop, wrtv=TRUE, para=15) # TRUE
## End(Not run)
```

isCOP.permsym

Is a Copula Permutation Symmetric

Description

Numerically set a logical whether a copula is *symmetric* (Nelsen, 2006, p. 38), or has *exchangeable* variables, or is *permutation symmetric* (Joe, 2014, p. 66) (refer also to *radial symmetric*, [isCOP.radsym](#)). A copula $\mathbf{C}(u, v)$ is permutation symmetric if and only if for any $\{u, v\} \in [0, 1]$ the following holds

$$\mathbf{C}(u, v) = \mathbf{C}(v, u).$$

The computation is (can be) CPU intensive and refer also to the extensive documentation and computational examples within [isCOP.radsym](#) (secs. **Note** and **Examples**) and the conceptual contributions of the *L-comoments* ([lcomCOP](#)) to symmetry.

Usage

```
isCOP.permsym(cop=NULL, para=NULL, delta=0.005, tol=1e-4, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters, if needed, to pass to the copula;
delta	The increment of $\{u, v\} \mapsto [0 + \Delta\delta, 1 - \Delta\delta, \Delta\delta]$;
tol	A tolerance on the check for symmetry, default 1 part in 10,000, which is the test for the $\equiv 0$ (zero equivalence, see source code); and
...	Additional arguments to pass to the copula.

Value

A logical TRUE or FALSE is returned.

Author(s)

W.H. Asquith

References

- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
 Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[LzCOPpermsym](#), [isCOP.radsym](#)

Examples

```
## Not run:
isCOP.permsym(cop=GHcop, para=1.3) # TRUE
## End(Not run)
```

isCOP.PQD

The Positively Quadrant Dependency State of a Copula

Description

Numerically determine the global property of the *positively quadrant dependency* (PQD) characteristic of a copula as described by Nelsen (2006, p. 188). The random variables X and Y are PQD if for all (x, y) in $\mathcal{R}^2$ when $H(x, y) \geq F(x)G(y)$ for all (x, y) in $\mathcal{R}^2$ and thus by the copula $\mathbf{C}(u, v) \geq uv$ for all (u, v) in $\mathcal{I}^2$. Alternatively, this means that $\mathbf{C}(u, v) \geq \mathbf{\Pi}$, and thus it can be said that it is globally “greater” than independence ($uv = \mathbf{\Pi}$; [P](#)).

Nelsen (2006) shows that a copula is PQD when

$$0 \leq \beta_{\mathbf{C}}, 0 \leq \gamma_{\mathbf{C}}, \text{ and } 0 \leq \rho_{\mathbf{C}} \leq 3\tau_{\mathbf{C}},$$

where $\beta_{\mathbf{C}}$, $\gamma_{\mathbf{C}}$, $\rho_{\mathbf{C}}$, and $\tau_{\mathbf{C}}$ are various copula measures of association or concordance that are respectively described in [blomCOP](#), [giniCOP](#), [rhoCOP](#), and [tauCOP](#). The concept of negatively quadrant

dependency (NQD) is the reverse: $C(u, v) \leq \Pi$ for all (u, v) in $\mathcal{I}^2$; so NQD is globally “smaller” than independence.

Conceptually, PQD is related to the probability that two random variables are simultaneously small (or simultaneously large) is at least as great as it would be if they were *independent*. The graph of a PQD copula lies on or above the copulatic surface of the *independence copula* Π , and conversely a NQD copula lies on or below Π .

Albeit a “global” property of a copula, there can be “local” variations in the PQD/NQD state. Points in $\mathcal{I}^2$ where $C(u, v) - \Pi \geq 0$ are locally PQD, whereas points in $\mathcal{I}^2$ where $C(u, v) - \Pi \leq 0$ are locally NQD. Lastly, readers are directed to the last examples in [wolfCOP](#) because as those examples involve the copulatic difference from independence $C(u, v) - \Pi = C(u, v) - \Pi$ with 3-D renderings.

Usage

```
isCOP.PQD(cop=NULL, para=NULL, uv=NULL, empirical=FALSE, verbose=TRUE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
uv	An optional R data.frame of U and V nonexceedance probabilities u and v for the random variables X and Y . This argument triggers different value return behavior (see Value);
empirical	A logical that will use sample versions for <i>Gini Gamma</i> , <i>Spearman Rho</i> , and <i>Kendall Tau</i> . This feature is <i>only</i> applicable if the copula is empirical and therefore the para argument is the data.frame of u and v , which will be passed along to sample version functions instead of copula (see Note);
verbose	A logical that will report the four concordance measures; and
...	Additional arguments to pass, which are then passed to subordinate functions.

Value

If $uv=NULL$ then a logical for the global property of PQD is returned but if argument uv is a data.frame, then an R list is returned, and that list holds the global condition in `global.PQD` and local condition assessments in `local.PQD` and `local.NQD`.

Note

The function `isCOP.PQD` will try brute force computations if subordinate calls to one or more functions fails. The user can use `...` to set the delta argument for [giniCOP](#), [rhoCOP](#), and (or) [tauCOP](#).

This function is not guaranteed to work using a *bivariate empirical copula* such as the following operation: `copPQD(cop=EMPIRcop, para=the.data)`. An evidently open problem for **copBasic** is how to support PQD assessment (either globally or locally) for empirical copulas. The τ_C for the bivariate empirical copula example `brute=TRUE|FALSE` to unity and γ_C and ρ_C reach maximum number of subdivisions on the numerical integration and thus fail. If an empirical bivariate copula is “Tau’d” to itself, is $\tau_C \equiv 1$ guaranteed? The τ_C computation relies on numerical partial derivatives

of the copula, whereas the γ_C and ρ_C use the copula itself. It seems in the end that use of sample versions of γ_C , ρ_C , and τ_C would be appropriate and leave the β_C as either copula or direct sample computation (see **Examples**).

SPECIAL DEMONSTRATION 1—Given the following,

```
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, para1=c(14.5), para2=c(1.45),
             alpha=0.51, beta=0.15, kappa=0.45, gamma=0.78)
D <- simCOP(n=500, cop=composite3COP, para=para, cex=0.5, col=1, pch=16)
```

the two different call types to isCOP.PQD for an empirical copula are illustrative:

```
global.only <- isCOP.PQD(cop=EMPIRcop, para=D, empirical=TRUE)
```

and

```
PQD.list <- isCOP.PQD(cop=EMPIRcop, para=D, empirical=TRUE, uv=D)
points(D, col=PQD.list$local.PQD+2, lwd=2) # red (if present) is local NQD
```

which in the former only returns the global PQD and the later returns an R list with global (global.PQD), local (local.PQD as well as local.NQD), and the four statistics (beta β_C , gamma γ_C , rho ρ_C , tau τ_C) used to determine global PQD.

SPECIAL DEMONSTRATION 1—Lastly, the ctype="bernstein" argument to the empirical copula can be used. Repeated iterations of the following will show that local quadrant dependency can appear slightly different when the bernstein argument is present. The simulation sample size is reduced considerably for this second example because of the CPU effort triggered by the *Bernstein extension* (see [EMPIRcop](#)) having been turned on.

```
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, para1=14.5, para2=1.45,
             alpha=0.51, beta=0.15, kappa=0.45, gamma=0.78)
D <- simCOP(n=50, cop=composite3COP, para=para, cex=0.5, col=1, pch=16)
PQD.A<- isCOP.PQD(cop=EMPIRcop, para=D, empirical=TRUE, uv=D)
points(D, col=PQD.A$local.PQD+2, lwd=2) # red (if present) is local NQD
PQD.B<- isCOP.PQD(cop=EMPIRcop, para=D, empirical=TRUE, uv=D, ctype="bernstein")
points(D, col=PQD.B$local.PQD+2, lwd=1, pch=3, cex=1.5)
```

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[blomCOP](#), [giniCOP](#), [rhoCOP](#), [tauCOP](#), [isCOP.LTD](#), [isCOP.RTI](#)

Examples

```
## Not run:
isCOP.PQD(cop=PSP) # TRUE
## End(Not run)

## Not run:
# Example concerning Empirical Bivariate Copula and sample versions for comparison.
set.seed(10); n <- 1000
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, para1=0.145, para2=1.45,
             alpha=0.81, beta=0.8)
D <- simCOP(n=n, cop=composite2COP, para=para, cex=0.5, col=rgb(0,0,0,0.2), pch=16)
#tauCOP(cop=EMPIRcop, para=D) # ??? but == 1
cor(D$U, D$V, method="kendall") # -0.3224705
blomCOP(cop=EMPIRcop, para=D) # -0.332
giniCOP(cop=EMPIRcop, para=D) # -0.3692037
GINI <- sum(abs(rank(D$U)+rank(D$V)-n-1)) - sum(abs(rank(D$U)-rank(D$V)))
print(GINI/as.integer(n^2/2)) # -0.369996
rhoCOP(cop=EMPIRcop, para=D) # ??? but fails
cor(D$U, D$V, method="spearman") # -0.456694
lmomco::lcomoms2(D)$T2 # 1.0000000 -0.4568357
# -0.4567859 1.0000000
## End(Not run)
```

isCOP.radsym

Is a Copula Radially Symmetric

Description

Numerically set a logical whether a copula is *radially symmetric* (Nelsen, 2006, p. 37) [*reflection symmetric*, Joe (2014, p. 64)] (refer also to *permutation symmetric*, [isCOP.permsym](#)). A copula $\mathbf{C}(u, v)$ is radially symmetric if and only if for any $\{u, v\} \in [0, 1]$ either of the following hold

$$\mathbf{C}(u, v) = u + v - 1 + \mathbf{C}(1 - u, 1 - v)$$

or

$$u + v - 1 + \mathbf{C}(1 - u, 1 - v) - \mathbf{C}(u, v) \equiv 0.$$

Thus, if the equality of the copula $\mathbf{C}(u, v) = \hat{\mathbf{C}}(u, v)$ (the *survival copula*), then radial symmetry exists: [COP](#) = [surCOP](#) or $\mathbf{C}(u, v) = \hat{\mathbf{C}}(1 - u, 1 - v)$. The computation is (can be) CPU intensive.

Usage

```
isCOP.radsym(cop=NULL, para=NULL, delta=0.005, tol=1e-4, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters, if needed, to pass to the copula;

delta	The increments of $\{u, v\} \mapsto [0 + \Delta\delta, 1 - \Delta\delta, \Delta\delta]$;
tol	A tolerance on the check for symmetry, default 1 part in 10,000, which is the test for the $\equiv 0$ (zero equivalence, see source code); and
...	Additional arguments to pass to the copula or derivative of a copula function.

Value

A logical TRUE or FALSE is returned.

Note

L-comoments and Radial and Permutation Symmetry — A natural question exists: *Is a radially symmetric copula characterized by the L-comoments for orders $r \geq 3$ as having values of zero?* Apparently, yes, and review of copula literature does not seem to directly address this question. Let us consider the two symmetrical copulas: the parameterless **PSP**(u, v) (see **PSP**) and the single parameter **PL**($u, v; \Theta$) (see **PLcop**) with the $\Theta_{\text{PL}} = 4.708664$ (**rhoCOP**). The two copulas have different radial symmetries as shown below.

```
J <- PLACKETTpar(rho=rhoCOP(cop=PSP)) # Spearman Rho = 0.4784176
isCOP.radsym(      cop=PSP)           # FALSE
isCOP.radsym(      cop=PLcop, para=J) # TRUE
UV <- simCOP(300, cop=PSP,  para=J, ploton=TRUE,  pch= 1, col="blue", seed=1)
UV <- simCOP(300, cop=PLcop, para=J, ploton=FALSE, pch=16, col="red" , seed=1)
```

Now, let us compute the L-comoments ($\tau_k^{[12][21]}$) (**lcomCOP**) from each copula in the code that follows. The L-correlations (Spearman Rho) are each about 0.478, which agree with the given ρ_C . The L-coskews (T3_PSP[12,21]) for the **PSP** are about -0.129 with those for the **PL** copula at zero. The **PSP** L-cokurtoses (T4_PSP[12,21]) are about 0.047, whereas those for the **PL** are zero. The [12], [21] equalities result from *permutation symmetry* (**isCOP.permsym**), but the nonzero L-cokurtoses for the **PSP** result from radial asymmetry.

```
lcmPSP <- lcomCOP(cop=PSP)
lcmPLA <- lcomCOP(cop=PLcop, para=J)
lcmPSP <- lapply(lcmPSP, function(i) round(i, digits=5))
lcmPLA <- lapply(lcmPLA, function(i) round(i, digits=5))
message( # L-correlations (T2), L-coskews (T3), L-cokurtoses (T4)
  "T2[12]_PSP == T2_PSP[21] =", lcmPSP$lcomUV[2], "=", lcmPSP$lcomVU[2], "\n",
  "T2[12]_PLA == T2_PLA[21] =", lcmPLA$lcomUV[2], "=", lcmPLA$lcomVU[2], "\n",
  "T3[12]_PSP == T3_PSP[21] =", lcmPSP$lcomUV[3], "=", lcmPSP$lcomVU[3], "\n",
  "T3[12]_PLA == T3_PLA[21] =", lcmPLA$lcomUV[3], "=", lcmPLA$lcomVU[3], "\n",
  "T4[12]_PSP == T4_PSP[21] =", lcmPSP$lcomUV[4], "=", lcmPSP$lcomVU[4], "\n",
  "T4[12]_PLA == T4_PLA[21] =", lcmPLA$lcomUV[4], "=", lcmPLA$lcomVU[4], "\n")
```

Now, **isCOP.radsym()** is false for all rotations (reflections) of the **PSP**, which requires that in magnitude that the L-coskews are equal but with reflections 3 and 4 rotating the couple to a $\rho_C = \tau_2^{[12]} = \tau_2^{[21]} = -0.478$. The L-cokurtoses are all equal with a change in sign with reflections 3 and 4. The **PSP** inherently has a bilateral symmetry about any diagonal, which makes the $r \geq 3$ L-comoments as mutual equals in magnitude.

```

lcmPSP1 <- lcomCOP(cop=COP, para=list(cop=PSP, reflect=1)) # default
lcmPSP2 <- lcomCOP(cop=COP, para=list(cop=PSP, reflect=2)) # 180 degrees
lcmPSP3 <- lcomCOP(cop=COP, para=list(cop=PSP, reflect=3)) # -90 degrees
lcmPSP4 <- lcomCOP(cop=COP, para=list(cop=PSP, reflect=4)) # +90 degrees
lcmPSP1 <- lapply(lcmPSP1, function(i) round(i, digits=5))
lcmPSP2 <- lapply(lcmPSP2, function(i) round(i, digits=5))
lcmPSP3 <- lapply(lcmPSP3, function(i) round(i, digits=5))
lcmPSP4 <- lapply(lcmPSP4, function(i) round(i, digits=5))
message( # L-coskews (T3), reflections 1~, 2~, 3~, 4~
  "1~T3[12] == T3[21] = ", lcmPSP1$lcomUV[3], " = ", lcmPSP1$lcomVU[3], "\n",
  "2~T3[12] == T3[21] = ", lcmPSP2$lcomUV[3], " = ", lcmPSP2$lcomVU[3], "\n",
  "3~T3[12] != T3[21] = ", lcmPSP3$lcomUV[3], " = ", lcmPSP3$lcomVU[3], "\n",
  "4~T3[12] != T3[21] = ", lcmPSP4$lcomUV[3], " = ", lcmPSP4$lcomVU[3], "\n")
# 1~T3[12] == T3[21] = -0.12949 = -0.12949
# 2~T3[12] == T3[21] = 0.12949 = 0.12949
# 3~T3[12] != T3[21] = 0.12949 = -0.12949
# 4~T3[12] != T3[21] = -0.12949 = 0.12949
message( # L-cokurtoses (T4), reflections 1~, 2~, 3~, 4~
  "1~T4[12] == T3[21] = ", lcmPSP1$lcomUV[4], " = ", lcmPSP1$lcomVU[4], "\n",
  "2~T4[12] == T3[21] = ", lcmPSP2$lcomUV[4], " = ", lcmPSP2$lcomVU[4], "\n",
  "3~T4[12] == T3[21] = ", lcmPSP3$lcomUV[4], " = ", lcmPSP3$lcomVU[4], "\n",
  "4~T4[12] == T3[21] = ", lcmPSP4$lcomUV[4], " = ", lcmPSP4$lcomVU[4], "\n")
# 1~T4[12] == T3[21] = 0.04677 = 0.04677
# 2~T4[12] == T3[21] = 0.04677 = 0.04677
# 3~T4[12] == T3[21] = -0.04677 = -0.04677
# 4~T4[12] == T3[21] = -0.04677 = -0.04677

```

Author(s)

W.H. Asquith

References

- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, Extremes in Nature—An approach using copulas: Springer, 289 p.

See Also

[isCOP.permsym](#), [lcomCOP](#)

Examples

```

# Radially symmetry is computationally intensive and relies on a gridded [0,1]x[0,1]
# space and laborious check on equality. Thus these examples are commented out for the
# R --timings check. Note, proof of radial symmetry absent of algebraic manipulation
# or verification is difficult and subject to the grid fineness to find a nonequality
# from which to immediately conclude FALSE. L-comoments could be used for simultaneous
# radial and permutation symmetry assessment and even beyond those by detection of
# no bilateral symmetry from any perspective (refer to the Note section above).

```

```

## Not run:
isCOP.radsym(cop=P) # TRUE

para <- list(cop1=PLcop, cop2=M, para1=c(.03), para2=NA, alpha=0.8, beta=0.5)
isCOP.radsym(      composite2COP, para=para) # FALSE
isCOP.permsym(      composite2COP, para=para) # FALSE
lcm <- lcomCOP(cop=composite2COP, para=para)
lcm <- lapply(lcm, function(i) round(i, digits=5))
message("      SpearmanRho, Lcoskew, Lcokurt\n",
        "Tk[12] ", paste(lcm$lcomUV[2:4], collapse=", "), "\n",
        "Tk[21] ", paste(lcm$lcomVU[2:4], collapse=", "))
#      SpearmanRho, Lcoskew, Lcokurt # Magitudes for L-comoment k>=3
# Tk[12] -0.25236, 0.22828, 0.01796 # are not equal because there is
# Tk[21] -0.25236, 0.17612, -0.05165 # no bilateral symmetry.
## End(Not run)

## Not run:
set.seed(120); theta <- 2
gh <- simCOP(n=34, cop=GHcop, para=theta, ploton=FALSE, points=FALSE) * 150
# Pretend gh is real data, the * 150 is to clearly get into an arbitrary unit system.

# The sort=FALSE is critical in the following two calls
fakeU <- lmomco::pp(gh[,1], sort=FALSE) # Weibull plotting position i/(n+1)
fakeV <- lmomco::pp(gh[,2], sort=FALSE) # Weibull plotting position i/(n+1)
uv <- data.frame(U=fakeU, V=fakeV); # our U-statistics

isCOP.radsym(cop=EMPIRCop, para=uv) # FALSE
isCOP.LTD(cop=EMPIRCop, para=uv) # TRUE
isCOP.RTI(cop=EMPIRCop, para=uv) # FALSE
isCOP.PQD(cop=EMPIRCop, para=uv, # TRUE
           empirical=TRUE) # need to set this true to avoid integrate()
# Blomqvist's Beta = 0.2941
#      Gini's Gamma = 0.5606
#      Spearman's Rho = 0.6584
#      Kendall's Tau = 0.5045

isCOP.radsym(cop=GHcop, para=theta) # FALSE
isCOP.LTD( cop=GHcop, para=theta) # TRUE
isCOP.RTI( cop=GHcop, para=theta) # TRUE
isCOP.PQD( cop=GHcop, para=theta) # TRUE
# Blomqvist's Beta = 0.5009
#      Gini's Gamma = 0.5591
#      Spearman's Rho = 0.6822
#      Kendall's Tau = 0.5000

# Notice that isCOP.RTI is not the same for empirical and theoretical.
# This shows the difficulty in tail dependence parameter estimation for
# small samples (refer also to Salvadori et al., 2007 p. 175).
## End(Not run)

```

Description

Numerically set a logical whether a copula is *right-tail increasing* (RTI) as described by Nelsen (2006, pp. 192–193) and Salvadori *et al.* (2007, p. 222). A copula $\mathbf{C}(u, v)$ is right-tail decreasing for RTI($V|U$) if and only if for any $v \in [0, 1]$,

$$\frac{\delta \mathbf{C}(u, v)}{\delta u} \leq \frac{v - \mathbf{C}(u, v)}{1 - u}$$

for almost all $u \in [0, 1]$. Similarly, a copula $\mathbf{C}(u, v)$ is right-tail decreasing for RTI($U|V$) if and only if for any $u \in [0, 1]$,

$$\frac{\delta \mathbf{C}(u, v)}{\delta v} \leq \frac{u - \mathbf{C}(u, v)}{1 - v}$$

for almost all $v \in [0, 1]$ where the later definition is controlled by the `wrtV=TRUE` argument.

The RTI concept is associated with the concept of *tail monotonicity* (Nelsen, 2006, p. 191). Specifically, but reference to Nelsen (2006) definitions and geometric interpretations is recommended, RTI($V|U$) (or RTI($V|U$)) means that the probability $P[Y > y | X > x]$ (or $P[X > x | Y > y]$) is a nondecreasing function of x (or y) for all y (or x).

A positive RTI of either RTI($V|U$) or RTI($U|V$) implies positively quadrant dependency (PQD, [isCOP.PQD](#)) but the condition of PQD does not imply RTI. Finally, the accuracy of the numerical assessment of the returned logical by `isCOP.RTI` is dependent on the the smallness of the `delta` argument passed into the function.

Usage

```
isCOP.RTI(cop=NULL, para=NULL, wrtV=FALSE, delta=0.005, ...)
```

Arguments

<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters, if needed, to pass to the copula;
<code>wrtV</code>	A logical to toggle between with respect to v or u (default);
<code>delta</code>	The increment of $\{u, v\} \mapsto [0 + \Delta\delta, 1 - \Delta\delta, \Delta\delta]$ set by <code>wrtV</code> ; and
<code>...</code>	Additional arguments to pass to the copula or derivative of a copula function.

Value

A logical TRUE or FALSE is returned.

Author(s)

W.H. Asquith

References

- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, Extremes in nature—An approach using copulas: Dordrecht, Netherlands, Springer, Water Science and Technology Library 56, 292 p.

See Also

[isCOP.LTD](#), [isCOP.PQD](#)

Examples

```
## Not run:
isCOP.RTI(cop=P, delta=0.01) # independence should be FALSE
# but function returns TRUE. Note, same logic for isCOP.LTD returns FALSE.
isCOP.RTI(cop=PSP) # TRUE : positive assoc.
isCOP.RTI(cop=PLACKETTcop, para=.15) # FALSE : negative assoc. Plackett
isCOP.RTI(cop=PLACKETTcop, para=15) # TRUE : positive assoc. Plackett
isCOP.RTI(cop=PLACKETTcop, wrtv=TRUE, para=.15) # FALSE : negative assoc. Plackett
isCOP.RTI(cop=PLACKETTcop, wrtv=TRUE, para=15) # TRUE : positive assoc. Plackett
## End(Not run)
```

isfuncCOP

Is a General Bivariate Function a Copula by Gridded Search?

Description

Is a general bivariate function a copula? Three properties are identified by Nelsen (2006, p. 10) for a bivariate copula $\mathbf{C}(u, v)$:

$$\mathbf{C}(u, 0) = 0 = \mathbf{C}(0, v) \quad \text{Nelsen 2.2.2a,}$$

$$\mathbf{C}(u, 1) = u \text{ and } \mathbf{C}(1, v) = v \quad \text{Nelsen 2.2.2b, and}$$

for every u_1, u_2, v_1, v_2 in $\mathcal{I}^2$ such that $u_1 \leq u_2$ and $v_1 \leq v_2$,

$$\mathbf{C}(u_2, v_2) - \mathbf{C}(u_2, v_1) - \mathbf{C}(u_1, v_2) + \mathbf{C}(u_1, v_1) \geq 0 \quad \text{Nelsen 2.2.2c.}$$

The last condition is known also as “2-increasing.” The `isfuncCOP` works along a gridded search in the domain $\mathcal{I}^2 = [0, 1] \times [0, 1]$ for the 2-increasing check with a resolution $\Delta u = \Delta v = \text{delta}$. Because there are plenty of true copula functions available in the literature it seems unlikely that this function provides much production utility in investigations. This function is provided because part of the objectives of the **copBasic** package is for instructional purposes. The computational overhead is far too great for relative benefit to somehow dispatch to this function all the time using the other copula utilities in this package.

Usage

```
isfuncCOP(cop=NULL, para=NULL, delta=0.002, ...)
```

Arguments

<code>cop</code>	A potential bivariate copula function that accepts two arguments for the u and v and parameters along argument <code>para</code> with option of additional arguments through the <code>...</code> argument;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>delta</code>	The $\Delta u = \Delta v$ of the grid edges; and
<code>...</code>	Additional arguments to pass to the copula function.

Value

A logical value of TRUE or FALSE is returned.

Author(s)

S. Kloibhofer (idea and most code) and W.H. Asquith (documentation)

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[densityCOP](#)

Examples

```
## Not run:
"NelsenEx2.11" <- function(u,v, ...) { # Nelsen (2006, exer. 2.11, p. 16)
  if(length(u) > 1 & length(v) > 1 & length(u) != length(v)) return(NA)
  if(length(u) == 1) u <- rep(u, length(v))
  if(length(v) == 1) v <- rep(v, length(u))
  return(sapply(1:length(u), function(i) { upv <- u[i] + v[i]
    if(2/3 <= upv & upv <= 4/3) return(min(c(u,v,1/3,upv-(2/3))))
    max(u[i]+v[i]-1, 0) })))
}
isfuncCOP(cop=NelsenEx2.11) # FALSE
## End(Not run)
```

JOcopB5

The Joe/B5 Copula (B5)

Description

The *Joe/B5 copula* (Joe, 2014, p. 170) is

$$\mathbf{C}_{\Theta}(u, v) = \mathbf{B5}(u, v) = 1 - \left((1-u)^{\Theta} + (1-v)^{\Theta} - (1-u)^{\Theta}(1-v)^{\Theta} \right)^{1/\Theta},$$

where $\Theta \in [1, \infty)$. The copula as $\Theta \rightarrow \infty$ limits to the *comonotonicity coupla* ($\mathbf{M}(u, v)$ and $\mathbf{M}$), as $\Theta \rightarrow 1^+$ limits to *independence copula* ($\mathbf{\Pi}(u, v)$; $\mathbf{P}$). Finally, the parameter Θ is readily computed from a *Kendall Tau* ([tauCOP](#)) by

$$\tau_{\mathbf{C}} = 1 + \frac{2}{2-\Theta} \left(\psi(2) - \psi(1+2/\Theta) \right),$$

where ψ is the digamma() function and as $\Theta \rightarrow 2$ then

$$\tau_{\mathbf{C}}(\Theta \rightarrow 2) = 1 - \psi'(2)$$

where ψ' is the trigamma() function.

Usage

```
JOcopB5(u, v, para=NULL, tau=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula;
<code>tau</code>	Optional Kendall Tau; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned. Otherwise if `tau` is given, then the Θ is computed and a list having

<code>para</code>	The parameter Θ , and
<code>tau</code>	Kendall Tau.

and if `para=NULL` and `tau=NULL`, then the values within `u` and `v` are used to compute Kendall Tau and then compute the parameter, and these are returned in the aforementioned list.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[M](#), [P](#)

Examples

```
# Upper tail dependency of Theta = pi --> 2 - 2^(1/pi) = 0.753131 (Joe, 2014, p. 171).
taildepCOP(cop=JOcopB5, para=pi)$lambdaU # 0.75313

# Blomqvist Beta of Theta = pi (Joe, 2014, p. 171).
blomCOP(cop=JOcopB5, para=pi)          # 0.5521328
3 - 4*(2*(1/2)^pi - (1/4)^pi)^(1/pi) # 0.5521328

## Not run:
# A test near the limiting Theta for trigamma()
UV  <- simCOP(cop=JOcopB5, para=2, n=10000)
para <- JOcopB5(UV[,1], UV[,2])$para
message("Tau difference ", round(2-para, digits=2), " is small.") #
## End(Not run)
```

```
## Not run:
# embedment of parameters for permutation asymmetry and rotation
para <- list(cop=JOcopB5, para=30, reflect=3, breve=0.5)
UV <- simCOP(cop=COP, para=list(cop=breveCOP, para=para), n=1000) #
## End(Not run)

## Not run:
# Performance of the parameter as function of Kendall tau to the sample
# Kendall tau. There will be some failures in the root solving as Tau
# goes towards 1.
ts <- seq(0,1, by=0.001); us <- runif(400) # Taus and a set pf random Us
ps <- sapply(ts, function(t) JOcopB5(tau=t)$para) # parameters from Taus
ps <- unlist(ps); names(ps) <- NULL # handling for solution failures
ps <- c(ps, rep(NA, length(ts)-length(ps))) # padding the solution failures
sts <- sapply(ps, function(p) { if(is.na(p)) return(NA) # sample Taus
  t <- cor(us, simCOPv(us, cop=JOcopB5, para=p), method="kendall")
  ifelse(is.null(t), return(NA), return(t)) })
plot(ts, sts, lwd=0.8, col=grey(0.6), las=1,
  xlab="Kendall tau", ylab="Sample Kendall tau")
abline(0,1, col="black", lwd=2) #
## End(Not run)
```

joeskewCOP

Joe's Nu-Skew and the copBasic Nu-Star of a Copula

Description

Compute the measure of *permutation asymmetry*, which can be thought of as *bivariate skewness*, named for the **copBasic** package as *Nu-Skew* ν_C of a copula according to Joe (2014, p. 66) by

$$\nu_C = 3E[UV^2 - U^2V] = 6 \int \int_{\mathcal{I}^2} (v - u) \mathbf{C}(u, v) du dv.$$

This definition is effectively the `type="nu"` for the function for which the multiplier 6 has been converted to 96 as explained in the **Note**.

Numerical results indicate $\nu_W \approx 0$ (**W**), $\nu_{\Pi} = 0$ (**P**), $\nu_M \approx 0$ (**M**), $\nu_{PL} \approx 0$ for all Θ (**PLcop**), and the $\nu_{GH}^* = 0$ (**GHcop**); copulas with mirror symmetry across the equal value line have $\nu_C = 0$.

Asymmetric copulas do exist. For example, consider an asymmetric Gumbel–Hougaard **GH** copula with $\Theta_p = (5, 0.8, p)$:

```
optimize(function(p) { nuskewCOP(cop=GHcop, para=c(5,0.8, p)) },
  c(0,0.99) )$minimum
UV <- simCOP(n=10000, cop=GHcop, c(5,0.8, 0.2836485)) # inspect the graphics
48*mean(UV$U*$V^2 - UV$U^2*UV$V) # -0.2847953 (not the 3rd parameter)
```

The minimization yields $\nu_{GH(5,0.8,0.2836485)} = -0.2796104$, which is close the expectation computed where $48 = 96/2$.

A complementary definition is supported, triggered by `type="nustar"`, and is computed by

$$\nu_{\mathbf{C}}^* = 12 \iint_{\mathcal{I}^2} (v + u) \mathbf{C}(u, v) \, du dv - 4,$$

which has been for the **copBasic** package, $\nu_{\mathbf{C}}^*$ is named as *Nu-Star*, which the 12 and the -4 have been chosen so that numerical results indicate $\nu_{\mathbf{W}}^* = -1$ (**W**), $\nu_{\mathbf{H}}^* = 0$ (**P**), and $\nu_{\mathbf{M}}^* = +1$ (**M**).

Lastly, the `uvlmoms` function provides for a quantile-based measure of bivariate skewness based on the difference $U - V$ that also is discussed by Joe (2014, p. 66).

Usage

```
joeskewCOP(cop=NULL, para=NULL, type=c("nu", "nustar", "nuskew"),
           as.sample=FALSE, brute=FALSE, delta=0.002, ...)

nuskewCOP(cop=NULL, para=NULL, as.sample=FALSE, brute=FALSE, delta=0.002, ...)
nustarCOP(cop=NULL, para=NULL, as.sample=FALSE, brute=FALSE, delta=0.002, ...)
```

Arguments

<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>type</code>	The type of metric to compute (nu and nuskew are synonymous for $\nu_{\mathbf{C}}$ and nustar is for $\nu_{\mathbf{C}}^*$);
<code>brute</code>	Should brute force be used instead of two nested <code>integrate()</code> functions to perform the double integration;
<code>delta</code>	The du and dv for the brute force integration using brute;
<code>as.sample</code>	A logical controlling whether an optional R data.frame in <code>para</code> is used to compute the sample $\hat{\nu}$ or $\hat{\nu}^*$ (see Note). If set to -1 , then the message concerning CPU effort will be suppressed; and
<code>...</code>	Additional arguments to pass.

Details

The implementation of `joeskewCOP` for **copBasic** provides the second metric of asymmetry, but why? Consider the results that follow:

```
joeskewCOP(cop=GHcop, para=c(5, 0.8, 0.2836485), type="nu")
# -0.2796104
joeskewCOP(cop=GHcop, para=c(5, 0.2836485, 0.8), type="nu")
# +0.2796103
joeskewCOP(cop=GHcop, para=c(5, 0.8, 0.2836485), type="nu")
# 0.3571276
joeskewCOP(cop=GHcop, para=c(5, 0.2836485, 0.8), type="nu")
# 0.3571279
tauCOP(cop=GHcop, para=c(5, 0.2836485, 0.8))
# 0.2443377
```

The demonstration shows—at least for the symmetry (switchability) of the 2nd and 3rd parameters (π_2 and π_3) of the asymmetric **GH** copula—that the first definition ν is magnitude symmetric but carries a sign change. The demonstration shows magnitude and sign stability for ν^* , and ends with *Kendall Tau* (**tauCOP**). Collectively, Kendall Tau (or the other *symmetric measures of association*, e.g. **blomCOP**, **footCOP**, **giniCOP**, **hoefCOP**, **rhoCOP**, **wolfCOP**) when combined with ν and ν^* might provide a framework for parameter optimization of the asymmetric **GH** copula (see below).

The asymmetric **GH**_(5,0.2836485,0.8) is not radial (**isCOP.radsym**) or permutation (**isCOP.permsym**), but if $\pi_2 = \pi_3$ then the resulting **GH** copula is not radially symmetric but is permutation symmetric:

```
isCOP.radsym( cop=GHcop, para=c(5, 0.2836485, 0.8)) # FALSE
isCOP.permsym(cop=GHcop, para=c(5, 0.2836485, 0.8)) # FALSE
isCOP.radsym( cop=GHcop, para=c(5, 0.8,      0.8)) # FALSE
isCOP.permsym(cop=GHcop, para=c(5, 0.8,      0.8)) # TRUE
```

The use of ν_C and ν_C^* with a *measure of association* is just suggested above for parameter optimization. Suppose we have **GH**_(5,0.5,0.7) with *Spearman Rho* $\rho = 0.4888$, $\nu = 0.001475$, and $\nu^* = 0.04223$, and the asymmetric **GH** couple is to be fit. Parameter estimation for the asymmetric **GH** is accomplished by

```
"fitGHcop" <- function(hats, assocfunc=rhoCOP, init=NA, eps=1E-4, ...) {
  H <- GHcop # shorthand for the copula
  "objfunc" <- function(par) {
    par[1]  <- ifelse(par[1] < 1, return(Inf), exp(par[1])) # edge check
    par[2:3] <- pnorm(par[2:3]) # detransform
    hp <- c(assocfunc(H, par), nuskewCOP(H, par), nustarCOP(H, par))
    return(sum((hats-hp)^2))
  }
  # Theta=1 and Pi2 = Pi3 = 1/2 # as default initial estimates
  if(is.na(init)) init <- c(1, rep(1/2, times=2))
  opt <- optim(init, objfunc, ...); par <- opt$par
  para <- c( exp(par[1]), pnorm(par[2:3]) )
  names(para) <- c("Theta", "Pi2", "Pi3")
  fit <- c(assocfunc(H, para), nuskewCOP(H, para), nustarCOP(H, para))
  txt <- c("AssocMeasure", "NuSkew", "NuStar")
  names(fit) <- txt; names(hats) <- txt
  if(opt$value > eps) warning("inspect the fit")
  return(list(para=para, fit=fit, given=hats, optim=opt))
}
father <- c(5,.5,.7)
densityCOPplot(cop=GHcop, para=father, contour.col=8)
fRho <- rhoCOP(  cop=GHcop, father)
fNu  <- nuskewCOP(cop=GHcop, father)
fStar <- nustarCOP(cop=GHcop, father)

child <- fitGHcop(c(fRho, fNu, fStar))$para
densityCOPplot(cop=GHcop, para=child, ploton=FALSE)

cRho <- rhoCOP(  cop=GHcop, child)
```

```

cNu    <- nuskeWcOP(cop=GHcop, child)
cStar <- nustarCOP(cop=GHcop, child)
message("Father stats: ", paste(fRho, fNu, fStar, sep=", "))
message("Child stats: ", paste(cRho, cNu, cStar, sep=", "))
message("Father para: ", paste(father, collapse=", "))
message("Child para: ", paste(child, collapse=", "))

```

The initial parameter estimate has the value $\Theta = 1$, which is *independence* for the one parameter **GH**. The two other parameters are set as $\pi_2 = \pi_3 = 1/2$ to be in the mid-point of their domain. The transformations using the $\log() \leftrightarrow \exp()$ and $\text{qnorm}() \leftrightarrow \text{pnorm}()$ functions in R are used to keep the optimization in the viable parameter domain. The results produce a fitted copula of **GH**_(4.907,0.5006,0.7014). This fit aligns well with the parent, and the three statistics are essentially matched during the fitting.

The ν_C^* can be similar to [rhoCOP](#), but differences do exist. In the presence of radial symmetry, ($\nu_C == 0$), the ν_C^* is nearly equal to *Spearman Rho* for some copulas. Let us test further:

```

p <- 10^seq(0,2,by=.01)
s <- sapply(p, function(t) nustarCOP(cop=GHcop, para=c(t)))
r <- sapply(p, function(t) rhoCOP(cop=GHcop, para=c(t)))
plot(p,s, log="x", type="l", col=3, lwd=3); lines(p,r)

```

Now let us add some asymmetry

```

s <- sapply(p, function(t) nustarCOP(cop=GHcop, para=c(t, 0.25, 0.75)))
r <- sapply(p, function(t) rhoCOP(cop=GHcop, para=c(t, 0.25, 0.75)))
plot(p,s, log="x", type="l", col=3, lwd=3); lines(p,r)

```

Now let us choose a different (the *Clayton*) copula

```

s <- sapply(p, function(t) nustarCOP(cop=CLcop, para=c(t)))
r <- sapply(p, function(t) rhoCOP(cop=CLcop, para=c(t)))
plot(p,s, log="x", type="l", col=3, lwd=3); lines(p,r)

```

Value

The value for ν_C or ν_C^* is returned.

Note

The ν_C definition is given with a multiplier of 6 on the integrals in order to agree with Joe (2014) relation that is also shown. However, in mutual parameter estimation experiments using a simple sum-of-square errors as shown in the **Details**, it is preferred to have ν_C measured on a larger scale. Where does the 96 then come from? It is heuristically made so that the upright and rotated cophalf (see **Examples** under [asCOP](#) and [bilmoms](#) for this copula) acquires ν_C values of +1 and -1, respectively. As a result to make back comparisons to Joe results, the ratios of 96 are made in this documentation.

The source code shows slightly different styles of division by the sample size as part of the sample estimation of the statistics. The $\hat{\nu}$ using just division by the sample size as testing indicates that this statistic is reasonably unbiased for simple copula. The $\hat{\nu}^*$ with similar division is a biased statistic

and the bias is not symmetrical in magnitude or sign it seems whether the $\hat{\nu}^*$ is positive or negative. The salient code is `spm <- ifelse(corsgn == -1, +2.4, +1.1)` within the sources for which the corrections were determined heuristically through simulation, and `corsgn` is the sign of the sample Spearman Rho through the `cor()` function of R.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[uvskew](#), [blomCOP](#), [footCOP](#), [giniCOP](#), [hoefCOP](#), [rhoCOP](#), [tauCOP](#), [wolfCOP](#)

Examples

```
nuskewCOP(cop=GHcop,para=c(1.43,1/2,1))*(6/96) # 0.005886 (Joe, 2014, p. 184; 0.0059)

## Not run:
joeskewCOP(          cop=GHcop, para=c(8, .7, .5)) # -0.1523491
joeskewCOP(          cop=GHcop, para=c(8, .5, .7)) # +0.1523472
# UV <- simCOP(n=1000, cop=GHcop, para=c(8, .7, .5)) # see the switch in
# UV <- simCOP(n=1000, cop=GHcop, para=c(8, .5, .7)) # curvature
## End(Not run)

## Not run:
para=c(19,0.3,0.8); set.seed(341)
nuskew <- nuskewCOP( cop=GHcop, para=para) # 0.3057744
UV <- simCOP(n=10000, cop=GHcop, para=para) # a large simulation
mean((UV$U - UV$V)^3)/(6/96)              # 0.3127398

# Two other definitions of skewness follow and are not numerically the same.
uvskew(u=UV$U, v=UV$V, umv=TRUE) # 0.3738987 (see documentation uvskew)
uvskew(u=UV$U, v=UV$V, umv=FALSE) # 0.3592739 ( or documentation uvlmoms)
# Yet another definition of skew, which requires large sample approximation
# using the L-comoments (3rd L-comoment is L-coskew).
lmomco::lmomoms2(UV)$T3 # L-coskew of the simulated values [1,2] and [2,1]
#           [,1]      [,2]
#[1,] 0.007398438 0.17076600
#[2,] -0.061060260 -0.00006613
# See the asymmetry in the two L-coskew values and consider this in light of
# the graphic produced by the simCOP() called for n=10,000. The T3[1,1] is
# the sampled L-skew (univariate) of the U margin and T3[2,2] is the same
# but for the V margin. Because the margins are uniform (ideally) then these
# for suitable large sample must be zero because the L-skew of the uniform
# distribution is by definition zero.
#
# Now let us check the sample estimator for sample of size n=300, and the
# t-test will (should) result in acceptance of the NULL hypothesis.
```

```

S <- replicate(60, nuskewCOP(para=simCOP(n=300, cop=GHcop, para=para,
                                       graphics=FALSE), as.sample=TRUE))

t.test(S, mu=nuskew)
# t = 0.004633, df = 59, p-value = 0.9963
# alternative hypothesis: true mean is not equal to 0.3057744
# 95 percent confidence interval:
#  0.2854282 0.3262150
# sample estimates:
# mean of x
# 0.3058216
## End(Not run)

## Not run:
# Let us run a large ensemble of copula properties that use the whole copula
# (not tail properties). We composite a Plackett with a Gumbel-Hougaard for
# which the over all association (correlation) sign is negative, but amongst
# these statistics with nuskew and nustar at the bottom, there are various
# quantities that can be extracted. These could be used for fitting.
set.seed(873)
para <- list(cop1=PLcop, cop2=GHcop, alpha=0.6, beta=0.9,
             para1=.005, para2=c(8.3,0.25,0.7))
UV <- simCOP(1000, cop=composite2COP, para=para) # just to show
blomCOP(composite2COP, para)      # -0.4078657
footCOP(composite2COP, para)      # -0.2854227
hoefCOP(composite2COP, para)      # +0.5713775
lcomCOP(composite2COP, para)$lcomUV[3] # +0.1816084
lcomCOP(composite2COP, para)$lcomVU[3] # +0.1279844
rhoCOP(composite2COP, para)       # -0.5688417
rhobevCOP(composite2COP, para)    # -0.2005210
tauCOP(composite2COP, para)       # -0.4514693
wolfCOP(composite2COP, para)      # +0.5691933
nustarCOP(composite2COP, para)    # -0.5172434
nuskewCOP(composite2COP, para)    # +0.0714987
## End(Not run)

```

joint.curvesCOP

Compute Coordinates of the Marginal Probabilities given joint AND or OR Probabilities

Description

Compute the coordinates of the *bivariate marginal probabilities* for variables U and V given selected probabilities levels t for a copula $\mathbf{C}(u, v)$ for v with respect to u . For the case of a **joint and** probability, symbolically the solution is

$$\Pr[U \leq v, V \leq v] = t = \mathbf{C}(u, v),$$

where $U \mapsto [t_i, u_j, u_{j+1}, \dots, 1; \Delta t]$ (an irregular sequence of u values from the i th value of t_i provided through to unity) and thus

$$t_i \mapsto \mathbf{C}(u = U, v),$$

and solving for the sequence of v . The index j is to indicate that a separate loop is involved and is distinct from i . The pairings $\{u(t_i), v(t_i)\}$ for each t are packaged as an R `data.frame`. This operation is very similar to the plotting capabilities in `level.curvesCOP` for *level curves* (Nelsen, 2006, pp. 12–13) but implemented in the function `joint.curvesCOP` for alternative utility.

For the case of a **joint or** probability, the *dual of a copula (function)* or $\tilde{\mathbf{C}}(u, v)$ (`duCOP`) from a copula (Nelsen, 2006, pp. 33–34) is used and symbolically the solution is:

$$\Pr[U \leq v \text{ or } V \leq v] = t = \tilde{\mathbf{C}}(u, v) = u + v - \mathbf{C}(u, v),$$

where $U \mapsto [0, u_j, u_{j+1}, \dots, t_i; \Delta t]$ (an irregular sequence of u values from zero through to the i th value of t) and thus

$$t_i \mapsto \tilde{\mathbf{C}}(u = U, v),$$

and solving for the sequence of v . The index j is to indicate that a separate loop is involved and is distinct from i . The pairings $\{u(t_i), v(t_i)\}$ for each t are packaged as an R `data.frame`.

Usage

```
joint.curvesCOP(cop=NULL, para=NULL, type=c("and", "or"),
               probs=c(0.5, 0.8, 0.90, 0.96, 0.98, 0.99, 0.995, 0.998),
               zero2small=TRUE, small=1E-6, divisor=100, delu=0.001, ...)
```

Arguments

<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>type</code>	What type of joint probability is to be computed;
<code>probs</code>	The joint probabilities t_i from which to compute the coordinates. The default values represent especially useful annual return period equivalents that are useful in hydrologic risk analyses;
<code>zero2small</code>	A logical controlling whether exactly zero value for probability are converted to a <code>small</code> value and exactly unity values for probability are converted to the value <code>1 - small</code> ; this logical is useful if transformation from probability space into standard normal variates or <i>Gumbel reduced variates</i> (refer to function <code>prob2grv()</code> in package lmomco) is later desired by the user for attendant graphics (see Examples section);
<code>small</code>	The value for <i>small</i> described for <code>zero2small</code> ;
<code>divisor</code>	A divisor on a computation of a Δt for incrementing through the irregularly-spaced u domain as part of the coordinate computation (refer to source code);
<code>delu</code>	A Δu for setup of the incrementing through the irregularly-space u domain as part of the coordinate computation (refer to source code); and
<code>...</code>	Additional arguments to pass to the <code>duCOP</code> function of copBasic or <code>uniroot()</code> function in R.

Value

An R list is returned with elements each of the given `probs`.

Note

The arguments `divisor` and `delu` provide flexibility to obtain sufficient smoothness in the coordinate curvatures for a given t . The pairings $\{u(t_i), v(t_i)\}$ for each t packaged as `data.frames` within the returned list each have their own unique length, and this is the reason that a single “master” `data.frame` is not returned by this function.

Extended Example—The code below shows both types of joint probability being computed using the default probs. The plotting is made in *Gumbel reduced variates* (GRV) (refer to `prob2grv` in package **lmomco**). The GRV transformation is somewhat suitable for the magnitude variation in and at tail depth of the probs. Also with the transformation is being used, the `zero2small` logical is kept at `TRUE`, which is appropriate. The `zero2small` being set is also useful if *standard normal variate* transformation (SNV) (by the `qnorm()` function in R) were used instead.

The *Gumbel–Hougaard* copula (`GHcop`) and a *reversed* Gumbel–Hougaard copula `rGH` are composited together by `composite2COP`. These were copula were chosen so that some asymmetry in the solutions by `joint.curvesCOP` could be seen. We begin by specifying symmetrical plotting limits for later use and then creating a function for the reversed Gumbel–Hougaard and setting the parameters and composition weights:

```
grvlim <- lmomco::prob2grv(c(0.25, 0.999)) # out to 1,000 years
"rGHcop" <- function(u,v, ...) { u + v - 1 + GHcop(1-u, 1-v, ...) }
para <- list(alpha=0.16, beta=0.67, cop1 =GHcop, cop2 =rGHcop,
              para1=4.5, para2=2.2)
tau    <- tauCOP(  cop=composite2COP, para=para) # Tau    = 0.351219
nuskew <- nuskewCOP( cop=composite2COP, para=para) # Nuskew = 0.084262
UV <- simCOP(n=1000, cop=composite2COP, para=para, snv=TRUE)
```

The code also computed the *Kendall Tau* (`tauCOP`) and *Nu-Skew* (`nuskewCOP`) and the results shown. The code finishes with a simulation by `simCOP` of the copula composition just for reference with axes in terms of SNV.

Next, we compute and plot the joint probability curves. The tolerance for the `uniroot` calls is reset from the R defaults because slight wooble in the numerical solutions exists otherwise. The `AND` and `OR` lists each provide `data.frames` from which successive graphic calls plot the lines. The second `plot()` call is commented out so that both sets of joint probability curves are drawn on the same plot.

```
AND <- joint.curvesCOP(cop=composite2COP, para=para, type="and",
                      divisor=1000, tol=.Machine$double.eps)
plot(grvlim, grvlim, type="n",
     xlab="GUMBEL REDUCED VARIATE IN U", ylab="GUMBEL REDUCED VARIATE IN V")
for(t in sort(as.numeric(names(AND)))) {
  UV <- get(as.character(t), AND)
  lines(lmomco::prob2grv(      UV$U), lmomco::prob2grv(      UV$V))
  text( lmomco::prob2grv(median(UV$U)), lmomco::prob2grv(median(UV$V)),
        as.character(round(1/(1-t)), digits=0))
}

OR <- joint.curvesCOP(cop=composite2COP, para=para, type="or",
                     divisor=1000, tol=.Machine$double.eps)
```

```

for(t in sort(as.numeric(names(OR)))) {
  UV <- get(as.character(t), OR)
  lines(lmomco::prob2grv(      UV$U), lmomco::prob2grv(      UV$V),
        col="red")
  text( lmomco::prob2grv(median(UV$U)), lmomco::prob2grv(median(UV$V)),
        as.character(round(1/(1-t))), digits=0), col="red")
}
mtext("Return Periods: black=cooperative risk, red=dual risk")
abline(0,1, col="grey50", lty=2) # dash line is simply and equal value line

```

Black Curves—The black curves represent the nonexceedance **joint and** condition. The black curves are a form of *level curve* ([level.curvesCOP](#)), but it seems appropriate to not name them as such because Nelsen’s examples and others usually have the level curves on an even step interval of probability such as 10-percent level curves. The complement of nonexceedance **joint and** is the probability level that either or both random variables (say “hazards”) U or V causes “failure” at the respective return period.

Red Curves—The red curves represent the nonexceedance **joint or** (inclusive) condition. The complement of nonexceedance **joint or** (inclusive) is the probability level that both random variables (say “hazards”) U or V must simultaneously (or dually) occur for “failure” at the respective return period. Note, there is obviously asymmetry in the **joint or** curves.

Interpretation—Because the two hazards can “cooperate” to cause failure ([coCOP](#)) for an equal level of protection (say 500-year event) relative to the complement of nonexceedance **joint or** (inclusive) condition ([surCOP](#)), the marginal probabilities must be considerably higher. Users are encouraged to review [copBasic-package](#) and the figure therein.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[diagCOPatf](#), [duCOP](#), [jointCOP](#), [joint.curvesCOP2](#), [level.curvesCOP](#)

Examples

See to the Note section

joint.curvesCOP2	Compute Coordinates of the Marginal Probabilities given joint AND or OR Probability
------------------	---

Description

Compute the coordinates of the bivariate marginal probabilities for variables U and V given selected probabilities levels t for a copula $\mathbf{C}(u, v)$ for u with respect to v . For the case of a **joint and** probability, symbolically the solution is

$$\Pr[U \leq v, V \leq v] = t = \mathbf{C}(u, v),$$

where $V \mapsto [t_i, t_j, t_{j+1}, \dots, 1; \Delta]$ (an irregular sequence of v values from the i th value of t_i provided through to unity) and thus

$$t_i \mapsto \mathbf{C}(u, v = V),$$

and solving for the sequence of u . The index j is to indicate that a separate loop is involved and is distinct from i . The pairings $\{u(t_i), v(t_i)\}$ for each t are packaged as an **R** data.frame. This operation is very similar to the plotting capabilities in [level.curvesCOP2](#) for *level curves* (Nelsen, 2006, pp. 12–13) but implemented in the function `joint.curvesCOP2` for alternative utility.

For the case of a **joint or** probability, the *dual of a copula (function)* or $\tilde{\mathbf{C}}(u, v)$ from a copula (Nelsen, 2006, pp. 33–34) is used and symbolically the solution is:

$$\Pr[U \leq v \text{ or } V \leq v] = t = \tilde{\mathbf{C}}(u, v) = u + v - \mathbf{C}(u, v),$$

where $V \mapsto [0, v_j, v_{j+1}, \dots, t_i; \Delta]$ (an irregular sequence of v values from zero through to the i th value of t) and thus

$$t_i \mapsto \tilde{\mathbf{C}}(u, v = V),$$

and solving for the sequence of u . The index j is to indicate that a separate loop is involved and is distinct from i . The pairings $\{u(t_i), v(t_i)\}$ for each t are packaged as an **R** data.frame.

Usage

```
joint.curvesCOP2(cop=NULL, para=NULL, type=c("and", "or"),
  probs=c(0.5, 0.8, 0.90, 0.96, 0.98, 0.99, 0.995, 0.998),
  zero2small=TRUE, small=1E-6, divisor=100, delv=0.001, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
type	What type of joint probability is to be computed;
probs	The joint probabilities for which to compute the coordinates. The default values represent especially useful annual return period equivalents that are useful in hydrologic risk analyses;

zero2small	A logical controlling whether precise zero value for probability are converted to a small value and precise unity values for probability are converted to the value $1 - \text{small}$; this logical is useful if transformation from probability space into standard normal variates or <i>Gumbel reduced variates</i> (GRV; refer to function <code>prob2grv()</code> in package lmomco) is later desired by the user for attendant graphics (refer to Examples section);
small	The value for <i>small</i> described for <code>zero2small</code> ;
divisor	A divisor on a computation of a Δ for incrementing through the v domain as part of the coordinate computation (refer to source code);
delv	A Δv for setup of the incrementing through the v domain as part of the coordinate computation (refer to source code); and
...	Additional arguments to pass to the <code>duCOP</code> function of copBasic or <code>uniroot()</code> function.

Value

An R list is returned with elements each of the given probs.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[diagCOPatf](#), [duCOP](#), [jointCOP](#), [joint.curvesCOP](#), [level.curvesCOP2](#)

Examples

```
# See also Note for joint.curvesCOP()
## Not run:
# Approach the joint curves from both "with respect two" perspectives---results same.
JCvwrtu <- joint.curvesCOP( cop=PSP, prob=0.98)$"0.98"
JCuwrtv <- joint.curvesCOP2(cop=PSP, prob=0.98)$"0.98"; lim <- c(2,5)
plot(qnorm(JCvwrtu$U),
     qnorm(JCvwrtu$V), type="l", lwd=6, col="grey50", xlim=lim, ylim=lim,
     xlab="STANDARD NORMAL VARIATE OF U", ylab="STANDARD NORMAL VARIATE OF V")
lines(qnorm(JCuwrtv$U), qnorm(JCuwrtv$V), col="red", lwd=2)
mtext("98th Joint Percentile Level Curve for PSP Copula") #
## End(Not run)
```

jointCOP

Compute Equal Marginal Probabilities Given a Single Joint AND or OR Probability for a Copula

Description

Given a single *joint probability* denoted as t for a copula $\mathbf{C}(u, v)$ numerically solve for bivariate marginal probabilities U and V such that they are also equal to each other ($u = v = w$). For the case of a **joint and** probability, the primary diagonal of the copula (Nelsen, 2006, pp. 12 and 16) is solved for by a simple dispatch to the [diagCOPatf](#) function instead. Symbolically the solution is

$$\Pr[U \leq v, V \leq v] = t = \mathbf{C}(w, w).$$

For the case of a **joint or** probability, the *dual of a copula (function)* or $\tilde{\mathbf{C}}(u, v)$ from a copula (Nelsen, 2006, pp. 33–34; [duCOP](#)) is used where symbolically the solution is

$$\Pr[U \leq v \text{ or } V \leq v] = t = \tilde{\mathbf{C}}(u, v) = u + v - \mathbf{C}(u, v),$$

or

$$\Pr[U \leq v \text{ or } V \leq v] = t = 2w - \mathbf{C}(w, w).$$

The function for type="or" tests $\tilde{\mathbf{C}}(0, 0)$ and if it returns NA or NaN then the lower limit for the rooting is treated as `.Machine$double.eps` instead of 0 (zero).

Usage

```
jointCOP(t, cop=NULL, para=NULL, type=c("and", "or"), ...)
```

Arguments

<code>t</code>	The joint probability level t ;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>type</code>	The type of joint probability is to be computed; and
<code>...</code>	Additional arguments to pass to the duCOP function of copBasic or <code>uniroot()</code> function.

Value

A vector of the equal u and v probabilities for the given type at the joint probability level of t . The vector includes the t as the third element.

Note

ENSEMBLE 1—Counting and Copula Probabilities from a Massive Sample Size: Simulations can be used to check and verify select copula concepts. Begin with a *Gumbel–Hougaard* copula $\mathbf{GH}(u, v) = \mathbf{C}_\Theta(u, v)$ having parameter $\Theta = 1.5$, which corresponds to a *Kendall Tau* $\tau_C = 1/3$ ([GHcop](#)). Next, simulate and count the number of either U or V exceeding the 99th percentile “event.” The event can occur either from the random variable U or from V with equal “loss.” If the event occurs in both U and V , the loss is just the same as if U or V occurred. So, if a design were at the 100-year level and thus a 0.01 chance of loss each year, then 500 losses in 50,000 years would be expected.

```
set.seed(89); n <- 50000
UV <- simCOP(n, cop=GHcop, para=1.5, graphics=FALSE)
length(UV$U[UV$U > 0.99 | UV$V > 0.99])      # 799 times (losses)
length(UV$U[UV$U > 0.99356 | UV$V > 0.99356]) # 500 times (losses)
```

Letting JP equal 0.99356, which forces the required acceptance of 500 losses for the design, has conditions of $UV\$U > \text{JP}$ **or** $UV\$V > \text{JP}$ as well as condition of $UV\$U > \text{JP}$ **and** $UV\$V > \text{JP}$. These three conditions are captured using the structure of the R code listed. Manual searching resulted in a value for JP equaling 0.99356, which produces the 500 count losses (acceptable losses). Thus, JP is a marginal bivariate probability (in this case equality between $U_{\text{crit.}} = V_{\text{crit.}}$ declared) necessary to attain a 99th percentile joint protection from loss. The magnitude for either U or V thus must exceed the 99th percentile, and this is what the code shows with 799 losses.

It is important to consider that unless U and V are in perfect positive correlation (e.g. $\mathbf{M}(u, v)$, [M](#), *Fréchet–Hoeffding upper-bound copula*), that protection from loss needs to be higher than 0.99 if the marginal risk is set at that level. Continuing, if the 99th percentile is the 100-year event, then design criteria should be about 155 years instead [`lmomco::prob2T(0.99356)`]. The user can readily see this with the switch to perfect independence with the $\mathbf{GH}(u, v)$ copula with $\Theta = 1$ and produce quite different results or extreme correlation with say $\Theta > 20$.

To provide the protection for 500 exceedances in $n = 50,000$ trials and for purposes of demonstration, balance the protection between U and V by setting their probabilities equal, then the theoretical joint probability is

```
diagCOPatf(0.99, 0.99, cop=GHcop, para=1.5)      # 0.9936887
jointCOP( 0.99,      cop=GHcop, para=1.5, type="and") # 0.9936887
```

and these two probabilities, which in reality are actually based on same computation (`diagCOPatf`), and nearly are the same as $\text{JP} = 0.99356$ that was determined by the manual searching on the simulated data.

A **mutually inclusive and** condition can be arranged as follows, and it is implicit in the definition that both loss events occur at the same time:

```
length(UV$U[UV$U > 0.99 & UV$V > 0.99])      # 208 losses ( simulated )
surCOP( 1-0.99, 1-0.99, cop=GHcop, para=1.5) * n # 209 losses (theoretical)
surfuncCOP(0.99, 0.99, cop=GHcop, para=1.5) * n # 209 losses (theoretical)
```

What are the expected number of exceedances if designs for U and V are built at $U = V = 0.99$ marginal probabilities for protection?

```

coCOP(1-0.99, 1-0.99, cop=GHcop, para=1.5) * n # 791 losses (theoretical)
# by the co-copula which from the copula as nonexceedances is
(1-COP( 0.99, 0.99, cop=GHcop, para=1.5)) * n # 791 losses (theoretical)

```

Note, the 791 losses is nearly equal to 799, but obviously not 500 as one might incorrectly have imagined strictly in a univariate world.

Now, a couple of questions can be asked about the **joint and** and **joint or** probabilities using the definition of a copula **COP** and the *dual of a copula (function)* ($\tilde{C}(u, v)$, **duCOP**), respectively:

```

# The AND nonexceedances:
length(UV$U[UV$U <= 0.99 & UV$V <= 0.99]) / n      # 0.98402   ( simulated )
COP(0.99, 0.99, cop=GHcop, para=1.5)                # 0.9841727 (theoretical)

# The OR nonexceedances:
length(UV$U[UV$U <= 0.99 | UV$V <= 0.99]) / n      # 0.99584   ( simulated )
duCOP(0.99, 0.99, cop=GHcop, para=1.5)             # 0.9958273 (theoretical)

```

How about inversion of $\tilde{C}(u, v)$ by jointCOP and check against the simulation?

```

jointCOP(0.99, cop=GHcop, para=1.5, type="or")[1]    # 0.9763951 ( theor. )
length(UV$U[UV$U <= 0.9763951 | UV$V <= 0.9763951])/n # 0.98982 ( simulated)

```

The second probability is a value quite near to 0.99. So, if one wants mutual loss protection, compute the inversion of the dual of a copula using jointCOP(..., type="or"). Let us say that 0.80 mutual loss protection is wanted

```

jointCOP(0.80, cop=GHcop, para=1.5, type="or")[1]    # 0.6561286
n - length(UV$U[UV$U <= 0.6561286 | UV$V <= 0.6561286])# 10049 losses ( sim.)
n - (1-0.8)*n                                         # 10000 losses (theoretical)

```

The example here shows numerical congruence of $10,049 \approx 10,000$. An opposing question is useful. How about a **mutually exclusive or** condition as nonexceedances?

```

# The mutually exclusive OR as nonexceedances:
length((UV$U[ (UV$U <= 0.99 | UV$V <= 0.99) &
! (UV$U <= 0.99 & UV$V <= 0.99)])) # 591 losses ( simulated )
# The mutually exclusive OR as exceedances:
length(UV$U[ (UV$U > 0.99 | UV$V > 0.99) &
! (UV$U > 0.99 & UV$V > 0.99)])) # 591 losses ( simulated )

```

It is clear that $208 + 591 = 799$ as shown earlier; notice how there are two ways to get at the 591 count. There are 208 mutual loss events and 591 occasions where either U or V is the causation of loss. For comparison, how many observed events by random variable were observed?

```

length(UV$U[ (UV$U > 0.99)]) # 519 ( simulated )
length(UV$U[ (UV$V > 0.99)]) # 491 ( simulated )

```

If the variables were perfectly uncorrelated ($\mathbf{P}(u, v)$, $\mathbf{P}$, *independence copula*) would be $519 + 491 = 1,007$ losses. But for the simulations here, 799 losses occurred, so $1,007 - 799 = 208$ losses were at the same time caused by U and V .

Both of the following lengths A and B are 799 and both represent a **joint or** condition—the operations do not represent a **mutually exclusive or** condition.

```
A <- length(UV$U[UV$U > 0.99]) + length(UV$U[UV$V > 0.99]) -
      length(UV$U[UV$U > 0.99 & UV$V > 0.99])
B <- length(UV$U[UV$U > 0.99 | UV$V > 0.99]) # A == B == 799
```

ENSEMBLE 2—Simulation Study using a Real-World Sample Size: Now with identities of sorts shown and described above, let us test a theoretically consistent version of a sample of size 150 repeated 1,000 times at the 98th percentile against loss by one or the other random variables or both for slightly correlated U or V again following the Gumbel–Hougaard copula. The losses incurred by mutual event occurrence is the same as if one or the other variables produced an event causing loss.

```
n <- 250; nsim <- 1000; EitherEvent <- 0.98; MutualEvent <- 0.99; Theta <- 1.5
PT <- jointCOP(EitherEvent, cop=GHcop, para=Theta, type="and")[1] # 0.9873537
DU <- jointCOP(MutualEvent, cop=GHcop, para=Theta, type="or" ) [1] # 0.9763951
```

This next code listing is a redundant example to the one that follows, but it is shown because a slight possibility of confusion in the vectorized conditional evaluations in R. This first example concretely changes the loss events into binary states and adds them up prior to the condition.

```
set.seed(894234)
EX1a <- sapply(1:nsim, function(i) {
  uv <- simCOP(n, cop=GHcop, para=Theta, graphics=FALSE)
  length(uv$U[ as.numeric(uv$U > PT) + as.numeric(uv$V > PT) >= 1 ]) })
t.test(EX1a, mu=(1-EitherEvent) * n)
```

The expected count $E[EX1a] = 5$ and the simulation run yielded 5.058. The `t.test()` function in R results in a statistically insignificant difference. The following two examples use a similar there. For sake of both code brevity and clarity, the examples here all restart the simulations at the expense of computation time.

```
set.seed(894234)
EX1b <- sapply(1:nsim, function(i) {
  uv <- simCOP(n, cop=GHcop, para=Theta, graphics=FALSE)
  length(uv$U[uv$U > PT | uv$V > PT]) })
t.test(EX1b, mu=(1-EitherEvent) * n)
```

The expected count $E[EX1b] = 5$ —the same results are shown as in the first example listing.

Next, let us demonstrate the dual of a copula for mutually occurring events.

```
set.seed(894234)
EX2 <- sapply(1:nsim, function(i) {
  uv <- simCOP(n, cop=GHcop, para=Theta, graphics=FALSE)
  length(uv$U[uv$U > DU & uv$V > DU]) })
t.test(EX2, mu=(1-MutualEvent) * n)
```

The expected count $E[EX2] = 2.5$ and the simulation run yielded 2.49. The `t.test()` again results in a statistically insignificant difference.

The $U = V = 0.9873537$ for the “and” and $U = V = 0.9763951$ for the “or” are not equal marginal probabilities. Taking the larger of the two marginal probabilities, the actual joint protection from mutual event occurrence can be computed:

```
duCOP(0.9873537, 0.9873537, cop=GHcop, para=Theta) # 0.9947075
(1-0.9947075)*n # which is about 1.32 events per 250 trials.
```

So, the larger protection in terms of joint probabilities provided by `EitherEvent` at 0.98 instead of `MutualEvent` at 0.99 with respective protection at the 0.9873537 provides a `MutualEvent` protection of 0.9947075.

```
set.seed(894234)
EX3 <- sapply(1:nsim, function(i) {
  uv <- simCOP(n, cop=GHcop, para=Theta, graphics=FALSE)
  length(uv$U[uv$U > 0.9873537 & uv$V > 0.9873537]) })
t.test(EX3, mu=(1-0.9947075) * n)
```

The expected count $E[EX3] = 1.32$ and the simulation run yielded 1.31. The `t.test()` again results in a statistically insignificant difference, and thus the result indistinguishable from the expectation.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[diagCOPatf](#), [duCOP](#), [joint.curvesCOP](#), [level.curvesCOP](#)

Examples

```
jointCOP(0.50, cop=GHcop, para=1.5, type="and") # 0.6461941 0.6461941 0.5000000
jointCOP(2/3, cop=GHcop, para=1.5, type="or" ) # 0.4994036 0.4994036 0.6666667

# See extended code listings and discussion in the Note section
```

Description

The *Joe–Ma copula* (Joe, 2014, p. 177), crediting Joe and Ma (2000), is

$$\mathbf{C}_{\Theta}(u, v) = \mathbf{JOMA}(u, v) = 1 - F_{\Gamma} \left(\left\{ [F_{\Gamma}^{(-1)}(1 - u; \Theta)]^{\Theta} + [F_{\Gamma}^{(-1)}(1 - v; \Theta)]^{\Theta} \right\}^{(1/\Theta)}; \Theta \right),$$

where $F_{\Gamma}(x; \Theta)$ is the *cumulative distribution function* of the *Gamma* distribution, $F_{\Gamma}^{(-1)}(f; \Theta)$ is the *quantile function* of the *Gamma* distribution, and $\Theta \in (-1, +1)$, which is a different parameter range than shown in Joe (2014, p. 177) because of numerical difficulties with the *Gamma* distribution and functional performance in context of **copBasic** design. The copula limits, as $\Theta \rightarrow -1$, to the *countermonotonicity coupla* (**W**(u, v); **W**), as $\Theta \rightarrow 1$, to the *comonotonicity copula* (**M**(u, v); **M**), and as $\Theta \rightarrow 0^{\pm}$, to the *independence copula* (**Π**(u, v); **P**), and as $\Theta \rightarrow +\infty$. The parameter Θ is readily computed from *Kendall Tau* (**tauCOP**) or *Spearman Rho* (**rhoCOP**) by root-solving methods. Because the formulation here uses a differing parameter range than shown in Joe (2014), integral formulations for Kendall Tau are not shown in this documentation.

Usage

```
JOMAcop(u, v, para=NULL, rhotau=NULL,
        cortype=c("kendall", "spearman", "tau", "rho"), ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula;
<code>rhotau</code>	Optional Kendall Tau or Spearman Rho and parameter <code>para</code> is returned depending on the setting of <code>cortype</code> . The <code>u</code> and <code>v</code> can be used for estimation of the parameter as computed through the setting of <code>cortype</code> ;
<code>cortype</code>	A character string controlling, if the parameter is not given, to use a Kendall Tau or Spearman Rho for estimation of the parameter. The name of this argument is reflective of an internal call to <code>stats::cor()</code> to the correlation (association) setting for Kendall Tau or Spearman Rho; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned. Otherwise if `tau` is given, then the Θ is computed and a list having

<code>para</code>	The parameter Θ , and
-------------------	------------------------------

tau Kendall Tau.

and if para=NULL and tau=NULL, then the values within u and v are used to compute Kendall Tau and then compute the parameter, and these are returned in the aforementioned list. Or if rho is given, then the Θ is computed and a similar list is returned having similar structure but with Spearman Rho instead.

Note

The implementation of the copula here makes use of the negative association part of Joe's parameterization and rotates the copula to positive association. Experiments show Gamma distribution limitations that are not symmetrical to the problem at hand as the shape parameter in the distribution gets large. The negative association part of the copula makes use of conversion to Joe (2014, p. 179) formula for the *lower bounds* of the copula. Internally, the parameter range is converted to $\Theta \in (-1, +1)$ and therefore $\Theta = 0$ is in the middle and independence. There are two thresholds for absolute value of the transformed Θ in the source code and these stem from estimation of where the Gamma distribution simply breaks down. The benefit of the parameterization for this copula is we get near complete comprehensiveness that is symmetrical in Kendall Tau or Spearman Rho, which does seem likely in the R source code that Joe (2014) has accompanying that book.

Remark, because reflection of $1 - v$ is used for the construction of the positive association, for $0 < \Theta \leq +1$, the copula is the copula of $(u, 1 - v)$; care will be needed in some situations to work with this fact. Also, though the formula above suggests *permutation symmetry* between (u, v) and (v, u) , simulations shown some tendency for skewing off the diagonal of the association. This is currently thought to be tied to the numerical derivative direction used in **copBasic** simulation; counter point is the following recipe:

```
UV <- simCOP(2000, cop=JOMAcop, para=-0.9, seed=1) # appears to
# "cry" !!!! towards the upper-right corner and this is the native
# formulation because the Theta is less than 0. Let us then have a
# check of permutation symmetry through all four rotations.
isCOP.permsym(cop=COP, para=list(cop=JOMAcop, para=-0.9, reflect=1))
# [1] TRUE
isCOP.permsym(cop=COP, para=list(cop=JOMAcop, para=-0.9, reflect=2))
# [1] TRUE
isCOP.permsym(cop=COP, para=list(cop=JOMAcop, para=-0.9, reflect=3))
# [1] FALSE
isCOP.permsym(cop=COP, para=list(cop=JOMAcop, para=-0.9, reflect=4))
# [1] FALSE
isCOP.permsym(cop=COP, para=list(cop=JOMAcop, para=+0.9, reflect=1))
# [1] FALSE
isCOP.permsym(cop=COP, para=list(cop=JOMAcop, para=+0.9, reflect=2))
# [1] FALSE
isCOP.permsym(cop=COP, para=list(cop=JOMAcop, para=+0.9, reflect=3))
# [1] TRUE
isCOP.permsym(cop=COP, para=list(cop=JOMAcop, para=+0.9, reflect=4))
# [1] TRUE
```

All this tells us is that when the principle direction of the points are along the primary diagonal from lower left to upper right that variables are not exchangeable and hence there appears some

directionality at the Gamma distribution itself within the what seems to be a symmetrical formula undoubtedly for the native form. Is this caused by the skewness itself of the Gamma? Will this be a source of L-coskewness of this copula implementation relative to *Gaussian* ([NORMcop](#)) or *Frank* ([FRcop](#)) copulas? Lastly, consult the following

```
lcomCOP(cop=JOMAcop, para=-0.6)
lcomCOP(cop=FRcop, para=FRcop(rhotau=tauCOP(cop=JOMAcop, para=-0.6))$para)
```

that shows us that the two copula for the same Kendall Tau have differing L-comoments of course but that the Frank as L-coskews of zero and the Joe–Ma does not. One more check is now made to provide more informative about the diagonal and symmetry or not:

```
d <- -0.9; t <- seq(0.001, 0.999, by=0.001)
UV <- simCOP(1000, cop=JOMAcop, para=d) # "crying" towards the upper right
abline(1, -1, col="red") # corner, so not symmetrical about the -1 diagonal.
u <- 0.5; der <- derCOP(cop=JOMAcop, u, t, para=-0.9, derdir="left")
plot(qnorm(t), qnorm(der), type="l", )
```

that tells us that a vertical second line at $u = c$ has a longer tail above the median and hence confirming that the Gamma distribution involvement must be producing the asymmetry.

Author(s)

W.H. Asquith

References

- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
- Joe, H., and Ma, C., 2000, Multivariate survival functions with a min-stable property: Journal of Multivariate Analyses, v. 75, no. 1, pp. 13–35, [doi:10.1006/jmva.1999.1891](#).

See Also

[M](#), [P](#), [W](#)

Examples

```
JOMAcop(rhotau=0.7, method="kendall")$para # 0.8645555

## Not run:
n <- 100 # Demonstrate "comprehensiveness" for the copula
d <- sort( sample(c(-1,1), n, replace=TRUE) * 10^runif(n, min=-0.5, max=0) )
for(i in 1:n) {
  UV <- simCOP(100, cop=JOMAcop, para=d[i]); mtext(d[i])
} #
## End(Not run)

## Not run:
# Close to maximum L-comoments and sample counterparts for general check
# against the postive and negative aspects and signs. It is interesting
```

```

# that there are some larger than anticipated differences with some of
# of the sample to the theory. Perhaps numerical improvements are needed?
UVp <- simCOP(100000, cop=JOMAcop, para=+0.6, graphics=FALSE, seed=1)
UVn <- simCOP(100000, cop=JOMAcop, para=-0.6, graphics=FALSE, seed=1)
#                               Lcomom:T3[12,21] Lcomom:T4[12,21]
lcomCOP(cop=JOMAcop, para=+0.6) #    0.014081858    0.014592670
#                               #    -0.014081858    0.014592670
lcomCOP(cop=JOMAcop, para=-0.6) #    0.014081858   -0.014592670
#                               #    0.014081858   -0.014592670
lmomco::lcomoms2(UVp, nmom=4) # $T3    0.0155001646    0.0161519394
#                               #   -0.0145945494    0.0148050507
lmomco::lcomoms2(UVn, nmom=4) # $T4    0.010777365    0.0143115158
#                               #   -0.0168424348   -0.0131075545 #
## End(Not run)

## Not run:
# A quick look at L-comoments of the copula with some remarks about limits.
# The numerical intergrations for the L-comoments "diverge" for
#   T3.12 NAs begin at -0.988, 0.987
#   T3.21 NAs begin at -0.988, -----
#   T4.12 NAs begin at -0.988, -----
#   T4.21 NAs begin at -0.988, -----
d <- seq(-1, 1, by=0.001)
t3.12 <- t4.12 <- rep(NA, length(d))
t3.21 <- t4.21 <- rep(NA, length(d))
for(i in seq_len(length(d))) {
  lcm <- lcomCOP(cop=JOMAcop, para=d[i])
  t3.12[i] <- lcm$lcomUV[3]; t4.12[i] <- lcm$lcomUV[4]
  t3.21[i] <- lcm$lcomVU[3]; t4.21[i] <- lcm$lcomVU[4]
}
plot(d, t3.12, col="skyblue4", type="l", lwd=3, ylim=c(-0.02, +0.02),
     xlab="JOMAP parameter", ylab="L-coskew and L-kurtosis")
lines(d, t4.12, col="skyblue4", lty=2, lwd=3)
lines(d, t3.21, col="red")
lines(d, t4.21, col="red", lty=2) #
## End(Not run)

```

Description

To begin, there are at least three terms in the literature for what appear as the same function supported by the `kfuncCOP` function. The *Kendall Function* also is known as *Kendall Distribution Function* (Nelsen, 2006, p. 163) and *Kendall Measure* (Salvadori *et al.*, 2007, p. 148). Each of these is dealt with in sequel to set the manner of the rather lengthy documentation for this function.

KENDALL FUNCTION—The *Kendall Function* (F_K) (Joe, 2014, pp. 419–422) is the cumulative distribution function (CDF) of the vector $\mathbf{U} = (U_1, U_2, \dots)$ or $\mathbf{U} = (u, v)$ (bivariate) where $\mathbf{U}$ is distributed as the copula: $\mathbf{U} \sim \mathbf{C}(u, v)$. Letting Z be the random variable for $\mathbf{C}(u, v)$: $Z = \mathbf{C}(u, v)$, the Kendall Function is defined as

$$F_K(z; \mathbf{C}) = \Pr[Z \leq z; \mathbf{U} \sim \mathbf{C}(u, v)],$$

where F_K is the nonexceedance probability of the joint probability z stemming from the $\mathbf{C}$. Note, unlike its univariate counterpart, $F_K(z)$ is rarely uniformly distributed (Nelsen *et al.*, 2001, p. 278). The inverse $F_K^{(-1)}(z)$ is implemented by the `kfuncCOPinv` function, which could be used for simulation of the correct joint probability using a single uniformly distributed $\sim U(0,1)$ random variable providing the nonexceedance probability of the Kendall function. A reminder is needed that Z is the **joint probability** and $F_K(z)$ is the Kendall Function.

Joe (2014) and others as cited list various special cases of $F_K(z)$, inequalities, and some useful identities suitable for validation study:

- For $\mathbf{M}(u, v)$ (see [M](#)): $F_K(z) = z$ for all $0 < z < 1$ for all $d \geq 2$ dimensions;
- For $\mathbf{W}(u, v)$ (see [W](#)): $F_K(z) = 1$ for all $0 < z < 1$ for $d = 2$ (bivariate only);
- For $\mathbf{\Pi}(u, v)$ (see [P](#)): $F_K(z) = z - z \log z$ for $0 < z < 1$ for $d = 2$ (bivariate only);
- For any $\mathbf{C}$: $z \leq F_K(z)$ for $0 < z < 1$; and
- For any $\mathbf{C}$: $E[Z] = 1 - \int_0^1 F_K(t) dt \geq z$ (Nelsen, 2001, p. 281) — Z expectation, not F_K !
- For any $\mathbf{C}$: $\tau_{\mathbf{C}} = 3 - 4 \int_0^1 F_K(t) dt$ (Nelsen, 2006, p. 163; see [tauCOP](#) [[Examples](#)]).
- For any $\mathbf{C}$: $F_K(t)$ does not uniquely determine the copula.

The last item is from Durante and Sempi (2015, p. 118), and later discussion herein will concern an example of theirs. By coincidence within a few days before receipt of the Durante and Sempi book, experiments using `kfuncCOP` suggested that numerically the Galambos ([GLcop](#)), Gumbel–Hougaard ([GHcop](#)), and Hüsler–Reiss ([HRcop](#)) extreme value copulas for the same *Kendall Tau* ($\tau_{\mathbf{C}}$) all have the same $F_K(t)$. Therefore, do all *EV*-copulas have the same Kendall Function? Well in fact, they do and Durante and Sempi (2015, p. 207) show that $F_K(z) = z - (1 - \tau_{\mathbf{C}})z \log(z)$ for an *EV*-copula.

Joe (2014, p. 420) also indicates that strength of *lower-tail dependence* ([taildepCOP](#)) affects $F_K(z)$ as $z \rightarrow 0^+$, whereas strength of *upper-tail dependence* affects $F_K(z)$ as $z \rightarrow 1^-$. (A demonstration of tail dependence dependence is made in section [Note](#).) Also compared to *comonotonicity copula* [[M](#)] there are no *countermonotonicity copula* ($\mathbf{W}_{d>2}$) for dimensions greater the bivariate (Joe, 2014, p. 214)

Joe (2014) does not explicitly list an expression of $F_K(z)$ that is computable directly for any $\mathbf{C}(u, v)$, and Nelsen (2006, p. 163) only lists a form (see later in documentation) for *Archimedean copulas*. Salvadori *et al.* (2007, eq. 3.47, p. 147) also list the Archimedean form; however, Salvadori *et al.* (2007, eq. 3.49, p. 148) **also list a form computable directly for any $\mathbf{C}(u, v)$** . Considerable numerical experiments and derivations involving the $\mathbf{\Pi}(u, v)$ copula and results for $K_{\mathbf{C}}(z)$ shown later, indicate that the correct Kendall form for any $\mathbf{C}(u, v)$ is

$$F_K(z) \equiv z + \int_z^1 \frac{\delta \mathbf{C}(u, t)}{\delta u} du,$$

where $t = \mathbf{C}^{(-1)}(u, z)$ for $0 \leq z \leq 1$, t can be computed by the `COPinv` function, and the partial derivative $\delta \mathbf{C}(u, t)/\delta u$ can be computed by the `derCOP` function. It is a curiosity that this form is not in Joe (2014), Nelsen *et al.* (2001, 2003), or Nelsen (2006), but just in Salvadori *et al.* (2007) of these books on copulas.

KENDALL MEASURE—The actual expression for any $\mathbf{C}(u, v)$ by Salvadori *et al.* (2007, eq. 3.49, p. 148) is for *Kendall Measure* ($K_{\mathbf{C}}$) of a copula:

$$K_{\mathbf{C}}(z) = z - \int_z^1 \frac{\delta \mathbf{C}(u, t)}{\delta u} du,$$

where $t = \mathbf{C}^{(-1)}(u, z)$ for $0 \leq z \leq 1$. Those authors report that $K_{\mathbf{C}}(z)$ is the CDF of a random variable Z whose distribution is $\mathbf{C}(u, v)$. This clearly appears to be the same meaning as Joe (2014) and Nelsen (2006). The minus “−” in the above equation is very important.

Salvadori *et al.* (2007, p. 148) report that “the function $K_{\mathbf{C}}(z)$ represents a fundamental tool for calculating the return period of extreme events.” The complement of $K_{\mathbf{C}}(z)$ is $\bar{K}_{\mathbf{C}}(z) = 1 - K_{\mathbf{C}}(z)$, and the $\bar{K}_{\mathbf{C}}(z)$ inverse

$$\frac{1}{1 - K_{\mathbf{C}}(z)} = \frac{1}{\bar{K}_{\mathbf{C}}(z)} = T_{\text{KC}}$$

is referred to as a *secondary return period* (Salvadori *et al.*, 2007, pp. 161–170).

KENDALL DISTRIBUTION FUNCTION—Nelsen (2006, p. 163) defines the *Kendall Distribution Function* (say $K_{\mathbf{C}}^*(t)$) as

$$K_{\mathbf{C}}^*(t) = t - \frac{\phi(t)}{\phi'(t^+)},$$

where $\phi(t)$ is a *generator function* of an *Archimedean copula* and $\phi'(t^+)$ is a one-sided derivative (Nelsen, 2006, p. 125), and $\phi(t)$ is $\phi(\mathbf{C}(u, v)) = \phi(u) + \phi(v)$. This same form also is listed by Salvadori *et al.* (2007, eq. 3.47).

Nelsen (2006) does not seem to list a more general definition for any $\mathbf{C}(u, v)$. Because there is considerable support for Archimedean copulas in **R**, **copBasic** has deliberately been kept from being yet another Archimedean-based package. This is made for more fundamental theory and pedagogic reasons without the algorithmic efficiency relative to the many convenient properties of Archimedean copulas.

The similarity of $F_K(z)$, $K_{\mathbf{C}}(z)$, and $K_{\mathbf{C}}^*(t)$, however, is obvious—research shows that there are no syntactic differences between $F_K(z)$ and $K_{\mathbf{C}}(t)$ and $K_{\mathbf{C}}^*(z)$ —they all are the CDF of the joint probability Z of the copula. Consider now that Salvadori *et al.* show $K_{\mathbf{C}}$ having the form $a - b$ and not a form $a + b$ as previously shown for $F_K(z)$. Which form is thus correct? The greater bulk of this documentation seeks to answer that question, and it must be concluded that Salvadori *et al.* (2007, eq. 3.49) definition for $K_{\mathbf{C}}(z)$ has a typesetting error.

Usage

```
kfuncCOP(z, cop=NULL, para=NULL, wrtV=FALSE, as.sample=FALSE,
          verbose=FALSE, subdivisions=100L,
          rel.tol=.Machine$double.eps^0.25, abs.tol=rel.tol, ...)
kmeasCOP(z, cop=NULL, para=NULL, wrtV=FALSE, as.sample=FALSE,
          verbose=FALSE, subdivisions=100L,
          rel.tol=.Machine$double.eps^0.25, abs.tol=rel.tol, ...)
```

Arguments

<code>z</code>	The values for z ;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>wrtV</code>	A logical to toggle between with respect to v or u (default);
<code>as.sample</code>	A control on whether an optional R data.frame in <code>para</code> is used to compute the empirical $\hat{F}_K(z)$. Let vector length of <code>para</code> be denoted m and $i = (1, \dots, m)$, if <code>as.sample=TRUE</code> , then for each of the probability pairs in <code>para</code> , the <i>empirical copula</i> (EMPIRCOP) function with additional arguments <code>...</code> is used to generate a vector $(0, F_{K,m}^\sharp, 1)$ of length $m + 2$, then a vector of $(0, z_m^\sharp, 1)$ again of length $m + 2$ where $z^\sharp = (i - 0.5)/m$ is created and linear interpolation by the <code>approx()</code> function in R for each of the z values in <code>z</code> is used to estimate $\hat{F}_K(z)$ (see source code). If <code>as.sample="genest"</code> , then the $\hat{F}_K(z)$ is estimated without interpolation using a simple empirical copula basis and “pseudo-observations” after Genest <i>et al.</i> (2006, p. 339). The default $\hat{F}_K$ (<code>as.sample=TRUE</code>) is the linear interpolation based on the default call of Weibull form of the empirical copula, but that can be confirmed by <code>ctype="weibull"</code> (see Note);
<code>verbose</code>	A logical suppressing warnings from <code>integrate()</code> in R that are usually related to “integral divergence” for $z \rightarrow 0^+$. The constructed behavior of <code>kfuncCOP</code> is to return $F_K(z \rightarrow 0^+) = 0$ if numerical integration returns NULL, and such a construction is made to avoid “end points not of opposite sign” during $F_K(z)$ inversion by kfuncCOPinv ;
<code>subdivisions</code>	Argument of same name passed to <code>integrate()</code> ,
<code>rel.tol</code>	Argument of same name passed to <code>integrate()</code> ,
<code>abs.tol</code>	Argument of same name passed to <code>integrate()</code> , and
<code>...</code>	Additional arguments to pass.

Value

The value(s) for $F_K(z)$ is returned.

Note

VALIDATION STUDY—A validation study using the *Independence copula* ($\Pi = uv$; **P**) with theoretical results of Joe (2014) and empiricism is readily performed using the expression for $F_K(z)$:

```
Z <- sort(c(0.01, seq(0,1, by=0.05), 0.99))      # ** Joint probabilities **
UV <- simCOP(n=4000, cop=P, graphics=FALSE);      kendF <- Z - Z*log(Z)
emp.kendF <- kfuncCOP(Z, para=UV, as.sample="genest") # emp. Kendall func
theo.kendF <- kfuncCOP(Z, cop=P) # theo. Kendall func, numeric integration
plot(Z, kendF, type="l", col="darkgreen", lwd=4, lty=2, xlim=c(0,1),
      xlab="COPULA(u,v) VALUE [JOINT PROBABILITY]", ylim=c(0,1),
      ylab="KENDALL FUNCTION, AS NONEXCEEDANCE PROBABILITY") # analytical
points(Z, kendF, col="green", lwd=1, lty=2, pch=16) # theoretical values
points(Z, emp.kendF, col="blue", lwd=2, cex=1.5)    # empirical values
```

```

lines( Z, theo.kendF, col="red") # theoretical line by numerical integration
lines( Z, kfuncCOP(Z, cop=M), lty=4) # dash-dot line (perfect correlation)
mtext("Kendall Functions: Independence Copula")

```

The figure produced shows that the theoretical relation in Joe (2014) is correct because the empirical values from the simulated sample (*Empirical Kendall Function*; Nelsen *et al.*, 2003) match other curves quite well. Rerunning the above code with either **M** (**M**) or **W** (**W**) copulas will show that the special cases listed above are consistent with the empirical estimates. The case of **W**(u, v) is degenerate at $z = 0$; so, the empirical computation is in error for the smallest z given because of interpolation. The **M** copula has F_K along the equal value line (1:1) line.

Now, consider a more comprehensive demonstration using the **N4212cop** copula with some relatively weak dependence in $\Theta = 1.17$.

```

Theta <- 1.17; print(rhoCOP(cop=N4212cop, para=Theta)) # Spearman Rho = 6/10
Z <- sort(c(0.01, seq(0,1, by=0.05), 0.99)) # ** Joint probabilities **
UV <- simCOP(n=16000, cop=N4212cop, para=Theta, graphics=TRUE)
empir.kendF <- kfuncCOP(Z, as.sample=TRUE, para=UV, ctype="weibull")
kwrtU <- kfuncCOP(Z, cop=N4212cop, para=Theta, wrtV=FALSE)
kwrtV <- kfuncCOP(Z, cop=N4212cop, para=Theta, wrtV=TRUE)
plot(Z, empir.kendF, type="p", col="red", lwd=7, lty=2, xlim=c(0,1),
     xlab="COPULA(u,v) VALUE [JOINT PROBABILITY]", ylim=c(0,1),
     ylab="KENDALL FUNCTION, AS NONEXCEEDANCE PROBABILITY")
abline(0,1, lty=2, lwd=0.8); mtext("Kendall Functions: N4212(1.17) Copula")
lines(Z, kwrtU, col="skyblue4", lwd=4, lty=2)
lines(Z, kwrtV, col="cyan2", lwd=2, lty=2)

```

The figure produced again shows congruence between the two theoretical computations and the empirical curve. Now, consider another comprehensive demonstration using the **PLACKETTcop** copula with some strong negative dependence in $\Theta = 0.04$.

```

Theta <- 0.04
Z <- sort(c(0.01, seq(0,1, by=0.05), 0.99)) # ** Joint probabilities **
UV <- simCOP(n=2600, cop=PLACKETTcop, para=Theta, graphics=TRUE)
empir.kendF <- kfuncCOP(Z, as.sample="hazen", para=UV)
kwrtU <- kfuncCOP(Z, cop=PLACKETTcop, para=Theta, wrtV=FALSE)
kwrtV <- kfuncCOP(Z, cop=PLACKETTcop, para=Theta, wrtV=TRUE)
plot(Z, empir.kendF, type="p", col="red", lwd=7, lty=2, xlim=c(0,1),
     xlab="COPULA(u,v) VALUE [JOINT PROBABILITY]", ylim=c(0,1),
     ylab="KENDALL FUNCTION, AS NONEXCEEDANCE PROBABILITY")
abline(0,1, lty=2, lwd=0.8); mtext("Kendall Function: Plackett Copula")
lines(Z, kwrtU, col="skyblue4", lwd=4, lty=2)
lines(Z, kwrtV, col="cyan2", lwd=1, lty=2)

```

The figure so produced again shows congruence between the two theoretical computations and the empirical curve.

Another comparison of $F_K(z)$ is useful and concerns *lower-* and *upper-tail dependency* parameters (**taildepCOP**) with a comparison of three different copula all having the same Kendall Tau. The following code computes and draws the respective $F_K(z)$:

```
# Given a Kendall Tau of 0.4 and the GHcop, N4212, and Plackett copulas
# parameters respectively are:
Phi <- 1.666667; Nu <- 1.111119; Mu <- 6.60344
Z <- seq(0.005, 0.995, by=0.005) # ** Joint probabilities **
GHkenf <- kfuncCOP(Z, cop=GHcop, para=Phi, wrtV=FALSE)
N4212kenf <- kfuncCOP(Z, cop=N4212cop, para=Nu, wrtV=FALSE)
PLkenf <- kfuncCOP(Z, cop=PLACKETTcop, para=Mu, wrtV=FALSE)
plot(qnorm(GHkenf), Z, type="l", col="black", lwd=2,
     xlab="KENDALL FUNCTION, AS STANDARD NORMAL VARIATES", xlim=c(-3,3),
     ylab="COPULA(u,v) VALUE, AS NONEXCEEDANCE PROBABILITY", ylim=c(0,1))
lines(qnorm(N4212kenf), Z, col="red", lwd=2)
lines(qnorm(PLkenf), Z, col="blue", lwd=2)
```

The red curve for the **N4212**($\Theta=1.1$) copula (**N4212cop**) is higher on the left, which shows the impact of its larger lower-tail dependency ($\lambda_{\text{GH}}^L=0 < \lambda_{\text{N4212}}^L=0.535$), whereas the black curve for the **GH**($\Theta=1.67$) copula (**GHcop**) is similarly (about same magnitude) higher on the right, which shows the impact of its larger upper-tail dependency ($\lambda_{\text{N4212}}^L=0 < \lambda_{\text{GH}}^U=0.484$). The blue curve for the **PL**($\Theta = 6.60$) copula (**PLACKETTcop**) nearly overwrites on the left the **GH** curve, which is a reflection of both copulae having zero lower-tail dependencies ($\lambda_{\text{GH}}^L = \lambda_{\text{PL}}^L = 0$). Finally, as anticipated by λ^U , the curve on the right for the **N4212** is just slightly larger for the **PL** because the $\lambda_{\text{PL}}^U=0 < \lambda_{\text{N4212}}^U=0.134$ (a small difference however), and again on the right, the **N4212** curve is considerably smaller than the **GH** because $\lambda_{\text{N4212}}^U=0 < \lambda_{\text{GH}}^U=0.484$.

Durante and Sempi (2015, p. 118) provide an example of two copula (C_1 and C_2) having the same $F_K(z) = \min(2z, 1)$. Let us check that out:

```
"C1" <- function(u,v, ...) {
  if (length(u) == 1) { u <- rep(u, length(v)) } else
  if (length(v) == 1) { v <- rep(v, length(u)) }
  sapply(1:length(u), function(i) {
    min(c(u[i], max(c(v[i]/2, u[i]+v[i]-1)))) })
}
"C2" <- function(u,v, ...) {
  if (length(u) == 1) { u <- rep(u, length(v)) } else
  if (length(v) == 1) { v <- rep(v, length(u)) }
  sapply(1:length(u), function(i) { g <- 1/2
    max(c(0, u[i]+v[i]-1, min(c(u[i], v[i]-g)), min(c(u[i]-g, v[i])))) })
}
DSkf <- function(t) sapply(t, function(z) min(c(2*z, 1)))
zs <- seq(0,1, by=.01); plot(zs, DSkf(zs), col="red", cex=3) # theory
lines(zs, kfuncCOP(zs, cop=C1), lwd=4, col="lightgreen")
lines(zs, kfuncCOP(zs, cop=C2), lwd=1, col="black")
```

The plot produced shows indeed that the numerical operations by `kfuncCOP` solidly work on these two strictly singular copulas C_1 and C_2 that are different from the two singular **M** and **W** copulas. The $F_K(z)$ curves exactly matching the theoretical curve provided by Durante and Sempi are produced.

CONVERSATIONAL ASIDE—Interestingly, Durante and Sempi (2015, pp. 115–121), like other authors of major works cited herein, do not list a general expression for the $F_K(z)$ as a function

of any $C(u, v)$. Those authors well develop the idea of Kendall Function and show results, but for the author (Asquith), the jump based of Theorem 3.9.2 to an expression, such as shown above for $F_K(z)$ based on $C(u, t)/\delta u$, as an usable form for any $C(u, v)$ is difficult.

This is a fascinating topic because, if the Kendall Function is a reasonably important component of copula theory, then why so much difficulty in finding a canonical reference? For this one piece, whereas so much of **copBasic** features are quite notationally clear in say Nelsen (2006) or Joe (2014) but somehow not for the Kendall Function.

Perhaps to the professional mathematicians, the descriptions (nomenclature) used in all but Salvadori *et al.* (2007) are clear to intended readers. But even Salvadori *et al.* seemingly show theirs in error—perhaps the author (Asquith) has missed something fundamental, but the validations shown in this documentation prove at least that kfuncCOP does what it is supposed to be doing but perhaps for the wrong reasoning. Lastly, Durante and Sempi (2015) have an error in their expression for Kendall Tau as a function of $F_K(z)$ (see [tauCOP](#)).

SECONDARY RETURN PERIOD—As for “Kendall Measure” (after the lead of Salvadori *et al.* [2007, pp. 161–170]), the following code shows for purposes of discussing *secondary return period* that $F_K(z)$ is correct and once and for all as defined in this documentation $F_K(z) \neq K_C(z)$. The secondary return period (T_K) is the expected interarrival time between events exceeding a T -year joint probability either from U , from V , or both. Let us use the 100-year level (primary return period; $T = 100$), the Gumbel–Hougaard copula ([GHcop](#)) $F_K^{\text{GH}(3.055)}(z)$ where the choice of $\Theta = 3.055$ is made to match discussion in [copBasic-package](#) and Salvadori *et al.* (2007, table 3.3, p. 166).

```
set.seed(1)
# Gumbel-Hougaard [Kendall Tau=0.67267 (Salvadori et al. [2007])]
Tyear <- 100; ANEP <- 1 - 1/Tyear; nsim <- 30000; ix <- seq_len(nsim)
1 / (1 - kfuncCOP(ANEP, cop=GHcop, para=3.055) ) # 148.28 years
BarT <- sapply(1:30, function(i) {
  UV <- simCOP(n=nsim, cop=GHcop, para=3.055, graphics=FALSE)
  nsim/sum(GHcop(UV$U, UV$V, para=3.055) > ANEP) })
message("# BarBarT = ", round(mean(BarT), digits=2),
        " years and StdDev(BarT) = ", round(sd(BarT), digits=2), " years")
# BarBarT = 151.78 years and StdDev(BarT) = 9.74 years
```

The mean of some 30 replications of a large sample simulation run for $F_K^{\text{GH}}(\Theta=3.055)_K(z=0.99)$ demonstrates empirical results that closely approximate theory $151.78 \approx 148.28$, and thus congruence is shown that the definition for $F_K(z)$ must be correct and that for $K_C(z)$ is incorrect. Table 3.3 in Salvadori *et al.* (2007, table 3.3) lists the secondary return period as 148.3 years, which matches the output of kfuncCOP and empirical results shown here.

Some additional details on secondary return period from Salvadori *et al.* (2007). Letting t_* be some critical joint exceedance probability level, $\bar{F}_K(z)$ be the complement of $F_K(z)$, those authors (p. 166) name *super-critical events* having $\bar{F}_K(t_*) < 1 - t_*$. Thus, the secondary return period $(\bar{F}_K(t_*)^{-1} = T_K)$ must be greater than the primary return period (t_*^{-1}) , which is the case here ($T_K=148.3 > T=100$) (Salvadori *et al.*, 2007, p. 166).

Salvadori *et al.* (2007) provide considerable discussion of T_K and some highlights are:

- (p. 162) events equally or exceeding probability $F_K(1 - 1/t_*)$ or having return intervals $\geq T_K$ “represent [a] class of potentially dangerous events, the *outliers*, and [F_K can be used to] introduce an *ad hoc* return period for such destructive events.”

- (p. 162) “primary return period $[T]$... only takes into account the fact that a prescribed critical event is expected to appear once in a given time interval $[T]$... $\bar{F}_K(t_*)$ provides the *exact probability* that a potentially destructive event will happen at any given realization of Z ... and T_K gives the expected time for such an outlier to occur.”
- (p. 164) “[T] only predicts that a critical event is *expected* to appear once in a given time interval ... would be more important to be able to calculate (1) the probability that a super-critical [sic.] event will occur at any given realization of $[Z]$, and (2) how long it takes, *on average*, for a super-critical event to appear.”
- (p. 164) “the function $\bar{F}_K$ turns the difficult analysis of bivariate dynamics of X and Y into a simpler one-dimensional problem.”

Author(s)

W.H. Asquith

Source

The comprehensive demonstrations are shown in the **Note** because of a sign convention and (or) probability convention incompatibility with the equation shown by Salvadori *et al.* (2007, p. 148). Initial source code development for **copBasic** was based on an understanding that the terms the “Kendall Function” and “Kendall Measure” might somehow have separate meanings—that the author must be blamed for misunderstanding the requisite nomenclature—this is evidently not true.

The $K_C(z)$ as shown herein simply can not reproduce $F_K(z; \Pi) = z - z \log z$ for the Π copula unless the “ $-$ ” sign in the $K_C(z)$ equation is changed to a “ $+$ ” to become the $F_K(z)$ definition as shown. The investigative effort to determine a valid function kmeasCOP was further complicated by fact that neither Durante and Sempi (2015), Joe (2014), Nelsen (2006), and others do not actually present a general equation for $F_K(z)$ computation for any $C(u, v)$.

Because of the subtle differences evidently between “Kendall functions” (lower case), an explicit derivation for $F_K(z; \Pi)$ is informative to confirm what is meant by the *Kendall Function* as defined by $F_K(z)$. Starting with $z = \Pi(u, v) = uv$, then

$$v(z) = t = \Pi^{(-1)}(u, z) = z/u, \text{ and}$$

$$\frac{\delta}{\delta u} \Pi(u, t) = \frac{\delta}{\delta u} uv \rightarrow v = \frac{z}{u},$$

substitution can now proceed:

$$F_K(z; \Pi) = z + \int_z^1 \frac{\delta}{\delta u} \Pi(u, t) du = z + \int_z^1 \frac{z}{u} du,$$

which simplifies to

$$F_K(z; \Pi) = z + [z \log(u)] \Big|_z^1 = z + z[\log(1) - \log(z)] = z - z \log z,$$

which matches the special case shown by Joe (2014) for the independence copula (Π ; **P**). It is obvious thus that the “ $+$ ” is needed in the $F_K(z)$ definition in order to stay consistent with the basic form and integration limits shown by Salvadori *et al.* (2007) for $K_C(z)$.

Lastly, an example under **W** demonstrates the span of the Kendall function based on **M** and **W** having permutation asymmetry inserted by **breveCOP**. The demonstration shows that the Kendall function is a distribution unto itself with its own nonexceedance probabilities.

References

- Durante, F., and Sempi, C., 2015, Principles of copula theory: Boca Raton, CRC Press, 315 p.
- Genest, C., Quessy, J.F., Rémillard, B., 2006, Goodness-of-fit procedures for copula models based on the probability integral transformation: Scandinavian Journal of Statistics, v. 33, no. 2, pp. 337–366.
- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Nelsen, R.B., Quesada-Molina, J.J., Rodríguez-Lallena, J.A., Úbeda-Flores, M., 2001, Distribution functions of copulas—A class of bivariate probability integral transforms: Statistics and Probability Letters, v. 54, no. 3, pp. 277–282.
- Nelsen, R.B., Quesada-Molina, J.J., Rodríguez-Lallena, J.A., Úbeda-Flores, M., 2003, Kendall distribution functions: Statistics and Probability Letters, v. 65, no. 3, pp. 263–268.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, Extremes in nature—An approach using copulas: Dordrecht, Netherlands, Springer, Water Science and Technology Library 56, 292 p.

See Also

[kfuncCOPinv](#), [tauCOP](#), [derCOP](#), [derCOP2](#), [derCOPinv](#), [derCOPinv2](#)

Examples

```
## Not run:
# Salvadori et al. (2007, p. 148, fig. 3.5 [right])
zs <- c(0.0001, seq(0.01, 1, by=0.01), 0.9999)
plot(zs, kmeasCOP(zs, cop=GHCop, para=5), log="y", type="l", lwd=4,
      xlab="Z <= z", ylab="Kendall Function", xlim=c(0,1), ylim=c(0.001,1)) #
## End(Not run)
```

kfuncCOPinv

The Inverse Kendall Function of a Copula

Description

Compute the (numerical) inverse $F_K^{(-1)}(z) \equiv z(F_K)$ of the *Kendall Function* $F_K(z; \mathbf{C})$ ([kfuncCOP](#)) of a copula $\mathbf{C}(u, v)$ given nonexceedance probability F_K . The z is the joint probability of the random variables U and V coupled to each other through the copula $\mathbf{C}(u, v)$ and the nonexceedance probability of the probability z is F_K —statements such as “probabilities of probabilities” are rhetorically complex so pursuit of word precision is made herein.

Usage

```
kfuncCOPinv(f, cop=NULL, para=NULL, subdivisions=100L,
            rel.tol=.Machine$double.eps^0.25, abs.tol=rel.tol, ...)
```

Arguments

f	Nonexceedance probability ($0 \leq F_K \leq 1$);
cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
subdivisions	Argument of same name passed to <code>integrate()</code> through kfuncCOP ,
rel.tol	Argument of same name passed to <code>integrate()</code> through kfuncCOP ,
abs.tol	Argument of same name passed to <code>integrate()</code> through kfuncCOP , and
...	Additional arguments to pass.

Value

The value(s) for $z(F_K)$ are returned.

Note

The L-moments of Kendall Functions appear to be unresearched. Therefore, the [kfuncCOPlmom](#) and [kfuncCOPlmoms](#) functions were written. These compute L-moments on the CDF $F_K(z)$ and not the quantile function $z(F_K)$ and thus are much faster than trying to use [kfuncCOPinv](#) in the more common definitions of L-moments. A demonstration of the mean (first L-moment) of the Kendall Function numerical computation follows:

```
# First approach
"afunc" <- function(f) kfuncCOPinv(f, cop=GHcop, para=pi)
integrate(afunc, 0, 1) # 0.4204238 with absolute error < 2.5e-05
# Second approach
kfuncCOPlmom(1, cop=GHcop, para=pi) # 0.4204222
```

where the first approach uses $z(F_K)$, whereas the second method uses integration for the mean on $F_K(z)$.

Author(s)

W.H. Asquith

References

- Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.
- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[kfuncCOP](#)

Examples

```
## Not run:
Z <- c(0,0.25,0.50,0.75,1) # Joint probabilities of a N4212cop
kfuncCOPinv(kfuncCOP(Z, cop=N4212cop, para=4.3), cop=N4212cop, para=4.3)
# [1] 0.0000000 0.2499984 0.5000224 0.7500112 1.0000000
## End(Not run)
```

kfuncCOPlmoms

The L-moments of the Kendall Function of a Copula

Description

Compute the L-moments of the *Kendall Function* ($F_K(z; \mathbf{C})$) of a copula $\mathbf{C}(u, v)$ where the z is the joint probability of the $\mathbf{C}(u, v)$. The *Kendall Function* (or *Kendall Distribution Function*) is the cumulative distribution function (CDF) of the joint probability Z of the coupla. The expected value of the $z(F_K)$ (mean, first L-moment λ_1), because Z has nonzero probability for $0 \leq Z \leq \infty$, is

$$E[Z] = \lambda_1 = \int_0^{\infty} [1 - F_K(t)] dt = \int_0^1 [1 - F_K(t)] dt,$$

where for circumstances here $0 \leq Z \leq 1$. The ∞ is mentioned only because expectations of such CDFs are usually shown using $(0, \infty)$ limits, whereas integration of quantile functions (CDF inverses) use limits $(0, 1)$. Because the support of Z is $(0, 1)$, like the probability F_K , showing just it (∞) as the upper limit could be confusing—statements such as “probabilities of probabilities” are rhetorically complex. So, pursuit of word precision is made herein.

An expression for λ_r for $r \geq 2$ in terms of the $F_K(z)$ is

$$\lambda_r = \frac{1}{r} \sum_{j=0}^{r-2} (-1)^j \binom{r-2}{j} \binom{r}{j+1} \int_0^1 [F_K(t)]^{r-j-1} \times [1 - F_K(t)]^{j+1} dt,$$

where because of these circumstances the limits of integration are $(0, 1)$ and not $(-\infty, \infty)$ as in the usual definition of L-moments in terms of a distribution’s CDF. (Note, such expressions did not make it into Asquith (2011), which needs rectification if that monograph ever makes it to a 2nd edition.)

The mean, L-scale, coefficient of L-variation (τ_2 , LCV, L-scale/mean), L-skew (τ_3 , TAU3), L-kurtosis (τ_4 , TAU4), and τ_5 (TAU5) are computed. In usual nomenclature, the L-moments are $\lambda_1 = \text{mean}$, $\lambda_2 = \text{L-scale}$, $\lambda_3 = \text{third L-moment}$, $\lambda_4 = \text{fourth L-moment}$, and $\lambda_5 = \text{fifth L-moment}$, whereas the L-moment ratios are $\tau_2 = \lambda_2/\lambda_1 = \text{coefficient of L-variation}$, $\tau_3 = \lambda_3/\lambda_2 = \text{L-skew}$, $\tau_4 = \lambda_4/\lambda_2 = \text{L-kurtosis}$, and $\tau_5 = \lambda_5/\lambda_2 = \text{not named}$. It is common amongst practitioners to lump the L-moment ratios into the general term “L-moments” and remain inclusive of the L-moment ratios. For example, L-skew then is referred to as the 3rd L-moment when it technically is the 3rd L-moment ratio. There is no first L-moment ratio (meaningless); so, results from `kfuncCOPlmoms` function will canoncially show a NA in that slot. The coefficient of L-variation is τ_2 (subscript 2) and not *Kendall Tau* (τ). Sample L-moments are readily computed by several packages in R (e.g. **lmomco**, **lmom**, **Lmoments**, **POT**).

Usage

```
kfuncCOPlmom(r, cop=NULL, para=NULL, ...)

kfuncCOPlmoms(cop=NULL, para=NULL, nmom=5, begin.mom=1, ...)
```

Arguments

<code>r</code>	The r th order of a single L-moment to compute;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>nmom</code>	The number of L-moments to compute;
<code>begin.mom</code>	The r th order to begin the sequence <code>lambegr:nmom</code> for L-moment computation. The rarely used argument is means to bypass the computation of the mean if the user has an alternative method for the mean or other central tendency characterization in which case <code>begin.mom = 2</code> ; and
<code>...</code>	Additional arguments to pass.

Value

An R list is returned by `kfuncCOPlmoms` and only the scalar value of λ_r by `kfuncCOPlmom`.

<code>lambdas</code>	Vector of the L-moments. First element is λ_1 , second element is λ_2 , and so on;
<code>ratios</code>	Vector of the L-moment ratios. Second element is τ , third element is τ_3 and so on; and
<code>source</code>	An attribute identifying the computational source of the L-moments: “kfuncCOPlmoms”.

Note

The L-moments of Kendall Functions appear to be not yet fully researched. An interesting research direction would be the trajectories of the L-moments or *L-moment ratio diagrams* for the Kendall Function and the degree to which distinction between copulas becomes evident—such diagrams are in wide-spread use for distinguishing between univariate distributions. It is noted, however, that *Kendall Function L-moment ratio diagrams* might be of less utility than in the univariate world—recalling that a univariate distribution is uniquely characterized by its L-moments—because different copulas can have the same $F_K(z)$, such as all bivariate extreme value copulas (see also **Examples**).

```
Rhos <- c(0.001, 0.01, seq(0.05, 0.95, by=0.05), 0.99, 0.999)
L1 <- T2 <- T3 <- T4 <- Thetas <- vector(mode="numeric", length(Rhos))
for(i in 1:length(Thetas)) {
  Thetas[i] <- uniroot(function(p)
    Rhos[i] - rhoCOP(cop=PARETOcop, para=p), c(0,200))$root
  message("Rho = ", Rhos[i], " and Pareto theta = ",
    round(Thetas[i], digits=4))
  lmr <- kfuncCOPlmoms(cop=PARETOcop, para=Thetas[i], nmom=4)
  L1[i] <- lmr$lambdas[1]; T2[i] <- lmr$ratios[2]
```

```

      T3[i] <- lmr$ratios[3]; T4[i] <- lmr$ratios[4]
    }
    LMR <- data.frame(Rho=Rhos, Theta=Thetas, L1=L1, T2=T2, T3=T3, T4=T4)
    plot(LMR$Rho, LMR$T2, ylim=c(-0.04, 0.5), xlim=c(0, 1),
         xlab="Spearman Rho or coefficient of L-variation",
         ylab="L-moment ratio", type="l", col="black")
    lines(LMR$Rho, LMR$T3, lty=1, col="red"           )
    lines(LMR$Rho, LMR$T4, lty=1, col="green"         )
    lines(LMR$T2, LMR$T3, lty=2, col="blue"           )
    lines(LMR$T2, LMR$T4, lty=2, col="deepskyblue2")
    lines(LMR$T3, LMR$T4, lty=2, col="purple"         )

```

Author(s)

W.H. Asquith

References

Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.

See Also

[kfuncCOP](#)

Examples

```

## Not run:
kfuncCOPlmom(1, cop=P) # 0.5 * 0.5 = 0.25 is expected joint prob. of independence
#[1] 0.2499999 (in agreement with theory)

ThetaGH <- 4.21
Rho <- rhoCOP(cop=GHCop, para=ThetaGH)
ThetaHR <- uniroot(function(p) Rho - rhoCOP(cop=HRCop, para=p), c(0, 100))$root
ThetaHR <- uniroot(function(p) Rho - rhoCOP(cop=HRCop, para=p), c(0, 100))$root
ThetaGL <- uniroot(function(p) Rho - rhoCOP(cop=GLcop, para=p), c(0, 100))$root
ls.str(kfuncCOPlmoms(cop=GHCop, para=ThetaGH)) # Gumbel-Hougaard copula
# lambdas : num [1:5] 0.440617 0.169085 0.011228 -0.000797 0.000249
# ratios : num [1:5] NA 0.383750 0.066400 -0.004720 0.001470
#
# L-skew = 0.066400
ls.str(kfuncCOPlmoms(cop=HRCop, para=ThetaHR)) # Husler-Reiss copula
# lambdas : num [1:5] 0.439627 0.169052 0.011427 -0.000785 0.000249
# ratios : num [1:5] NA 0.384540 0.067590 -0.004640 0.001470
#
# L-skew = 0.067590
ls.str(kfuncCOPlmoms(cop=GLcop, para=ThetaGL)) # Galambos copula
# lambdas : num [1:5] 0.440415 0.169079 0.011268 -0.000795 0.000248
# ratios : num [1:5] NA 0.383910 0.066650 -0.004700 0.001470
#
# L-skew = 0.066650
# These L-moments are extremely similar and within the numerics used.
# Extreme value copula all have the same Kendall Distribution function.
## End(Not run)

```

```
## Not run:
UV <- simCOP(200, cop=PLcop, para=1/pi, graphics=FALSE)
theta <- PLpar(UV[,1], UV[,2])
zs <- c(0.001, seq(0.01, 0.99, by=0.01), 0.999) # for later

# Take the sample estimated parameter and convert to joint probabilities Z
# Convert the Z to the Kendall Function estimates again with the sample parameter
Z <- PLcop(UV[,1], UV[,2], para=theta); KF <- kfuncCOP(Z, cop=PLcop, para=theta)

# Compute L-moments of the "Kendall function" and the sample versions
# and again see that the L-moment are for the distribution of the Z!
KNFlmr <- kfuncCOPlmoms(cop=PLcop, para=theta); SAMlmr <- lmomco::lmoms(Z)
knftxt <- paste0("Kendall L-moments: ",
                paste(round(KNFlmr$lambda, digits=4), collapse=", "))
samtxt <- paste0("Sample L-moments: ",
                paste(round(SAMlmr$lambda, digits=4), collapse=", "))

plot(Z, KF, xlim=c(0,1), ylim=c(0,1), col="black",
     xlab="COPULA(u,v) VALUE [JOINT PROBABILITY]",
     ylab="KENDALL DISTRIBUTION FUNCTION (KDF), AS NONEXCEEDANCE PROBABILITY")
rug(Z, side=1, col="red", lwd=0.5); rug(KF, side=2, col="red", lwd=0.5) # rug plots
lines(zs, kfuncCOP(zs, cop=PLcop, para=1/pi), col="darkgreen")
knf_meanZ <- KNFlmr$lambda[1]; sam_meanZ <- SAMlmr$lambda[1]
knf_mean <- kfuncCOP(knf_meanZ, cop=PLcop, para=theta) # theo. Kendall function
sam_mean <- kfuncCOP(sam_meanZ, cop=PLcop, para=theta) # sam. est. of Kendall func
points(knf_meanZ, knf_mean, pch=16, col="blue", cex=3)
points(sam_meanZ, sam_mean, pch=16, col="cyan", cex=2)
lines(zs, zs*zs*log(zs), lty=2, lwd=0.8) # dash ref line for independence
text(0.2, 0.30, knftxt, pos=4, cex=1); text(0.2, 0.25, samtxt, pos=4, cex=1)
text(0.2, 0.18, paste0("Notice uniform distribution of vertical axis rug.\n",
                      "A Critical remark with respect to to KDFs."), cex=1, pos=4)
legend("bottomright", c("Independence copula", "KDF of Plackett copula",
                      "Theoretical mean", "Sample mean"), bty="n", y.intersp=1.5,
      lwd=c(1, 1, NA, NA), lty=c(2, 1, NA, NA), pch=c(NA, NA, 16, 16),
      col=c("black", "darkgreen", "blue", "cyan"), pt.cex=c(NA, NA, 3, 2)) #
## End(Not run)
```

kfuncCOP_Pd

The Kendall (Distribution) Function of d-Dimensional Independence Copula

Description

Compute the *Kendall Distribution Function* of a *d*-Dimensional *independence copula* (P) (Joe, 2014, p. 420):

$$F_K(z; \mathbf{P}_d) = z + z \sum_{k=1}^{d-1} \left(-\log(k) \right)^k / k!,$$

where F_K is the nonexceedance probability of joint probability z stemming from $\mathbf{P}_d(u_1, 2, \dots, d)$.

Usage

```
kfuncCOP_Pd(z, d=2)
```

Arguments

<code>z</code>	The values for z ; and
<code>d</code>	Number of d dimensions.

Value

The value(s) for $F_K(z; \mathbf{P}_d)$ is returned.

Note

Though the **copBasic** is designed for bivariate relations, *vining* workflows could be used to produce d -dimensional relations from which Kendall functions could be empirically computed from d -variate simulations and a user might want an easy mechanism to look at *independence* on Kendall distribution function plots.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[kfuncCOP](#)

Examples

```
# The 0.50 joint probability has a 0.8465736 nonexceedance probability.
kfuncCOP_Pd(0.50, d=3) # 0.8465736

## Not run:
f <- seq(0,1, by=0.01)
plot(f, kfuncCOP(f, cop=P), type="l", lwd=7, col=grey(0.8),
      xlab="Kendall distribution function (joint probability)",
      ylab="Nonexceedance probability")
lines(f, kfuncCOP_Pd(f, d=2), lwd=1.5, col="black")
lines(f, kfuncCOP_Pd(f, d=3), col="seagreen3")
lines(f, kfuncCOP_Pd(f, d=4), col="seagreen4") #
## End(Not run)
```

kullCOP

Kullback–Leibler Divergence, Jeffrey Divergence, and Kullback–Leibler Sample Size

Description

Compute the *Kullback–Leibler Divergence*, *Jeffrey Divergence*, and *Kullback–Leibler sample size* following Joe (2014, pp. 234–237). Consider two densities $f = c_1(u, v; \Theta_f)$ and $g = c_2(u, v; \Theta_g)$ for two different bivariate copulas $C_1(\Theta_1)$ and $C_2(\Theta_2)$ having respective parameters Θ , then the Kullback–Leibler Divergence of f relative to g is

$$KL(f|g) = \int \int_{\mathcal{I}^2} g \log(g/f) \, du dv,$$

and Kullback–Leibler Divergence of g relative to f is

$$KL(g|f) = \int \int_{\mathcal{I}^2} f \log(f/g) \, du dv,$$

where the limits of integration $\mathcal{I}^2$ theoretically are closed on $[0, 1]^2$ but an open interval $(0, 1)^2$ might be needed for numerical integration. Note, in general $KL(f|g) \neq KL(g|f)$. The $KL(f|g)$ is the expected log-likelihood ratios of g to f when g is the true density (Joe, 2014, p. 234), whereas $KL(g|f)$ is the opposite.

This asymmetry leads to Jeffrey Divergence, which is defined as a symmetrized version of the two Kullback–Leibler Divergences, and is

$$J(f, g) = KL(f|g) + KL(g|f) = \int \int_{\mathcal{I}^2} (g - f) \log(g/f) \, du dv.$$

The variances of the Kullback–Leibler Divergences are defined as

$$\sigma_{KL(f|g)}^2 = \int \int_{\mathcal{I}^2} g [\log(g/f)]^2 \, du dv - [KL(f|g)]^2,$$

and

$$\sigma_{KL(g|f)}^2 = \int \int_{\mathcal{I}^2} f [\log(f/g)]^2 \, du dv - [KL(g|f)]^2.$$

For comparison of copula families f and g and taking an $\alpha = 0.05$, the Kullback–Leibler sample size is defined as

$$n_{fg} = [\Phi^{(-1)}(1 - \alpha) \times \eta_{KL}]^2,$$

where $\Phi^{(-1)}(t)$ is the quantile function for the standard normal distribution $\sim N(0,1)$ for nonexceedance probability t , and η_{KL} is the maximum of

$$\eta_{KL} = \max[\sigma_{KL(f|g)}/KL(f|g), \sigma_{KL(g|f)}/KL(g|f)].$$

The n_{fg} gives an indication of the sample size needed to distinguish f and g with a probability of at least $1 - \alpha = 1 - 0.05 = 0.95$ or 95 percent.

The **copBasic** features a naïve *Monte Carlo integration* scheme in the primary interface `kullCOP`, although the function `kullCOPint` provides for nested numerical integration. This later function is generally fast but suffers too much for general application from integral divergencies issued from the `integrate()` function in **R**—this must be judged in the light that the **copBasic** package focuses only on implementation of the function of the copula itself and numerical estimation of copula density (`densityCOP`) and not analytical copula densities or hybrid representations thereof. Sufficient “bread crumbs” are left among the code and documentation for users to re-implement if speed is paramount. Numerical comparison to the results of Joe (2014) (see **Examples**) suggests that the default Monte Carlo sample size should be more than sufficient for general inference with the expense of considerable CPU time; however, a couple of repeated calls of `kullCOP` would be advised and compute say the mean of the resulting sample sizes.

Usage

```
kullCOP(cop1=NULL, cop2=NULL, para1=NULL, para2=NULL, alpha=0.05,
        del=0, n=1E5, verbose=TRUE, sobol=FALSE, scrambling=0, ...)

kullCOPint(cop1=NULL, cop2=NULL, para1=NULL, para2=NULL, alpha=0.05,
           del=.Machine$double.eps^0.25, verbose=TRUE, ...)
```

Arguments

<code>cop1</code>	A copula function corresponding to copula f in Joe (2014);
<code>para1</code>	Vector of parameters or other data structure, if needed, to pass to the copula f ;
<code>cop2</code>	A copula function corresponding to copula g in Joe (2014);
<code>para2</code>	Vector of parameters or other data structure, if needed, to pass to the copula g ;
<code>alpha</code>	The α in the Kullback–Leibler sample size equation;
<code>del</code>	A small value used to denote the lo and hi values of the numerical integration: $lo = del$ and $hi = 1 - del$. If $del == 0$, then $lo = 0$ and $hi = 1$, which corresponds to the theoretical limits $\mathcal{I}^2 = [0, 1]^2$ and are defaulted here to $[0, 1]^2$ because the Monte Carlo algorithm is preferred for general application. The end point control, however, is maintained just in case pathological situations should arise;
<code>n</code>	<code>kullCOP</code> (Monte Carlo integration) only—the Monte Carlo integration simulation size;
<code>verbose</code>	A logical triggering a couple of status lines of output through the <code>message()</code> function in R ;
<code>sobol</code>	A logical triggering <i>Sobol sequences</i> for the Monte Carlo integration instead of the bivariate uniform distribution. The Sobol sequences are dependent on the randtoolbox package and the <code>sobol()</code> function of the randtoolbox package, and the Sobol sequences canvas the $\mathcal{I}^2$ domain for smaller n values than required if statistical independence is used for the Monte Carlo integration. Note, the randtoolbox at least at version 2.0.+ has “scrambling” of Sobol sequences temporarily disabled, and hence <code>scrambling=0</code> as default for <code>kullCOP</code> ;
<code>scrambling</code>	The argument of the same name for <code>randtoolbox::sobol</code> ; and
<code>...</code>	Additional arguments to pass to the <code>densityCOP</code> function.

Value

An R list is returned having the following components:

MonteCarlo.sim.size	kullCOP (Monte Carlo integration) only—The simulation size for numerical integration;
divergences	A vector of the Kullback–Leibler Divergences and their standard deviations: $KL(f g)$, $\sigma_{KL(f g)}$, $KL(g f)$, and $\sigma_{KL(g f)}$, respectively;
stdev.divergences	kullCOP (Monte Carlo integration) only—The standard deviation of the divergences and the variances;
Jeffrey.divergence	Jeffrey Divergence $J(f, g)$;
KL.sample.size	Kullback–Leibler sample size n_{fg} ; and
integrations	kullCOPint (numerical integration) only—An R list of the outer call of the <code>integrate()</code> function for the respective numerical integrals shown in this documentation.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[densityCOP](#), [vuongCOP](#)

Examples

```
# See another demonstration under the Note section of statTn().
## Not run:
# Joe (2014, p. 237, table 5.2)
# Gumbel-Hougaard and Plackett copulas below each have a Kendall Tau of about 0.5, and
# Joe (2014) lists in the table that Jeffrey Divergence is about 0.110 and Kullback-Leibler
# sample size is 133. Joe (2014) does not list the copula parameters just says Tau = 0.5.
# Joe (2014) likely did the numerical integrations using analytical solutions to probability
# densities and not rectangular approximations as in copBasic::densityCOP().
set.seed(1)
KL <- kullCOP(cop1=GHcop,      para1=2,
              cop2=PLACKETTcop, para2=11.40484, sobol=FALSE)
message("Jeffery Divergence is ",      round(KL$Jeffrey.divergence, digits=4),
        " and Kullback-Leibler sample size is ", KL$KL.sample.size, ".")
# Jeffery Divergence is 0.1106 and Kullback-Leibler sample size is 137.
set.seed(1)
KL <- kullCOP(cop1=GHcop,      para1=2,
              cop2=PLACKETTcop, para2=11.40484, sobol=TRUE )
```

```

message("Jeffery Divergence is ",          round(KL$Jeffrey.divergence, digits=4),
        " and Kullback-Leibler sample size is ", KL$KL.sample.size, ".")
# Jeffery Divergence is 0.3062 and Kullback-Leibler sample size is 136.

set.seed(1)
S <- replicate(20, kullCOP(cop1=GHcop, para1=2, cop2=PLACKETTcop, sobol=FALSE,
                           para2=11.40484, verbose=FALSE)$KL.sample.size)
print(as.integer(c(mean(S), sd(S)))) # 132 plus/minus 5
S <- replicate(2, kullCOP(cop1=GHcop, para1=2, cop2=PLACKETTcop, sobol=TRUE,
                           para2=11.40484, verbose=FALSE)$KL.sample.size)
# The two S in the later replication are both the same (136) for a sobol=TRUE
# does not produce variation and this is thought (June 2023) as a result
# of the disabled scrambling in the randtoolbox::sobol() function.
## End(Not run)

## Not run:
# Joe (2014, p. 237, table 5.3)
# Gumbel-Hougaard and Plackett copulas below each have a Spearman Rho of about 0.5, and
# Joe (2014) lists in the table that Jeffery Divergence is about 0.063 and Kullback-Leibler
# sample size is 210. Joe (2014) does not list the parameters and just says that Rho = 0.5.
# Joe (2014) likely did the numerical integrations using analytical solutions to probability
# densities and not rectangular approximations as in copBasic::densityCOP().
set.seed(1)
KL <- kullCOP(cop1=GHcop,          para1=1.541071,
               cop2=PLACKETTcop, para2=5.115658, sobol=FALSE)
message("Jeffery Divergence is ",          round(KL$Jeffrey.divergence, digits=4),
        " and Kullback-Leibler sample size is ", KL$KL.sample.size, ".")
# Jeffery Divergence is 0.0642 and Kullback-Leibler sample size is 213.
set.seed(1)
KL <- kullCOP(cop1=GHcop,          para1=1.541071,
               cop2=PLACKETTcop, para2=5.115658, sobol=TRUE )
message("Jeffery Divergence is ",          round(KL$Jeffrey.divergence, digits=4),
        " and Kullback-Leibler sample size is ", KL$KL.sample.size, ".")
# Jeffery Divergence is 0.2001 and Kullback-Leibler sample size is 206.

set.seed(1)
S <- replicate(20, kullCOP(cop1=GHcop, para1=1.541071, cop2=PLACKETTcop,
                           para2=5.115658, verbose=FALSE)$KL.sample.size)
print(as.integer(c(mean(S), sd(S)))) # 220 plus/minus 19
## End(Not run)

## Not run:
# Compare Jeffery Divergence estimates as functions of sample size when computed
# using Sobol sequences or not for Gumbel-Hougaard and Pareto copulas.
GHpar <- P Apar <- 2 # Spearman Rho = 0.6822339
Ns <- as.integer(10^c(seq(2.0, 3.5, by=0.01), seq(3.6, 5, by=0.05)))
JDuni <- sapply(1:length(Ns), function(i) {
  kullCOP(cop1=GHcop, para1=GHpar, verbose=FALSE,
           cop2=P A cop, para2=P Apar, n=Ns[i],
           sobol=FALSE)$Jeffrey.divergence })
JDsob <- sapply(1:length(Ns), function(i) {

```

```

      kullCOP(cop1=GHcop, para1=GHpar, verbose=FALSE,
              cop2=PAcop, para2=Papar, n=Ns[i],
              sobol=TRUE )$Jeffrey.divergence })
plot(Ns, JDuni, type="l", log="x", # black line, notice likely outliers too
      xlab="Simulation Sample Size", ylab="Jeffery Divergence")
lines(Ns, JDsob, col="red") # red line
legend("topright", c("Monte Carlo", "Sobol sequence"),
      lwd=c(1,1), col=c("black", "red"), bty="n")
print( c( mean(JDuni), sd(JDuni) ) ) # [1] 0.05915608 0.01284682
print( c( mean(JDsob), sd(JDsob) ) ) # [1] 0.07274190 0.01838939

# The developer notes that plotting KL.sample.size for sobol=TRUE shows
# what appears to be numerical blow up but the Jeffery Divergence does not.
KLuni <- sapply(1:length(Ns), function(i) {
      kullCOP(cop1=GHcop, para1=GHpar, verbose=FALSE,
              cop2=PAcop, para2=Papar, n=Ns[i],
              sobol=FALSE)$KL.sample.size })
KLsob <- sapply(1:length(Ns), function(i) {
      kullCOP(cop1=GHcop, para1=GHpar, verbose=FALSE,
              cop2=PAcop, para2=Papar, n=Ns[i],
              sobol=TRUE )$KL.sample.size })
plot(Ns, KLuni, type="l", log="xy", # black line, notice likely outliers too
      xlab="Simulation Sample Size", ylab="Kullback-Leibler Sample Size")
lines(Ns, KLSob, col="red") # red line
nideal <- kullCOPint(cop1=GHcop, para1=GHpar, cop2=PAcop, para2=Papar)$KL.sample.size
abline(h=nideal, col="green", lwd=3) # nideal sample size is about 210
legend("topright", c("Monte Carlo", "Sobol sequence", "Judged ideal sample size"),
      lwd=c(1,1,3), col=c("black", "red", "green"), bty="n")

# Let us attempt a visualization to highlight the differences in the two copula by
# simulation. First, using this n = nideal, being the apparent sample size to distinguish
# generally between the two copula having the same Spearman Rho. Do the segments help
# to visually highlight the differences? Next, ask would one judge the parents in the
# simulation being different knowing same Spearman Rho? (Note, the segments are all
# vertical because the U axis is the simulation and the V axis is the conditional
# probability given the U.)
set.seed(1); UVgh <- simCOP(nideal, GHcop, para=GHpar, graphics=FALSE)
set.seed(1); UVpa <- simCOP(nideal, PAcop, para=Papar, graphics=FALSE)
plot(c(0,1), c(0,1), type="n", xlab="U, nonexceedance probability",
      ylab="V, nonexceedance probability")
segments(UVgh[,1], UVgh[,2], x1=UVpa[,1], y1=UVpa[,2])
points(UVgh, col="lightgreen", pch=16, cex=0.8) # dots
points(UVpa, col="darkgreen", pch= 1, lwd=0.8) # circles
# Repeat the above n = nideal visualizations but with a change to n = nideal*10, and see
# then that there are visually striking shifts systematically in both both tails but also
# in the U in the interval (0.3, 0.7) belt but to a smaller degree than seen in the tails.
## End(Not run)

```

Description

Compute the *L-comoments* (Serfling and Xiao, 2007; Asquith, 2011) through the *bivariate L-moments (ratios)* ($\delta_{k;\mathbf{C}}^{[\dots]}$) of a copula $\mathbf{C}(u, v; \Theta)$. The L-comoments include *L-correlation* (*Spearman Rho*), *L-coskew*, and *L-cokurtosis*. As described by Brahimi *et al.* (2015), the first four bivariate L-moments $\delta_k^{[12]}$ for random variable $X^{(1)}$ or U with respect to (*wrt*) random variable $X^{(2)}$ or V are defined as

$$\delta_{1;\mathbf{C}}^{[12]} = 2 \int \int_{T^2} \mathbf{C}(u, v) \, du dv - \frac{1}{2},$$

$$\delta_{2;\mathbf{C}}^{[12]} = \int \int_{T^2} (12v - 6) \mathbf{C}(u, v) \, du dv - \frac{1}{2},$$

$$\delta_{3;\mathbf{C}}^{[12]} = \int \int_{T^2} (60v^2 - 60v + 12) \mathbf{C}(u, v) \, du dv - \frac{1}{2}, \text{ and}$$

$$\delta_{4;\mathbf{C}}^{[12]} = \int \int_{T^2} (280v^3 - 420v^2 + 180v - 20) \mathbf{C}(u, v) \, du dv - \frac{1}{2},$$

where the bivariate L-moments are related to the L-comoment ratios by

$$6\delta_k^{[12]} = \tau_{k+1}^{[12]} \quad \text{and} \quad 6\delta_k^{[21]} = \tau_{k+1}^{[21]},$$

where in otherwords, “the third bivariate L-moment $\delta_3^{[12]}$ is one sixth the L-cokurtosis $\tau_4^{[12]}$.” The first four bivariate L-moments yield the first five L-comoments. The terms and nomenclature are not easy and also the English grammar adjective “ratios” is not always consistent in the literature. The $\delta_{k;\mathbf{C}}^{[\dots]}$ are **ratios**. The sample L-comoments are supported by the **lmomco** package, and in particular for the bivariate case, they are supported by the `lcomoms2()` function of that package.

Similarly, the $\delta_k^{[21]}$ are computed by switching $u \rightarrow v$ in the polynomials within the above integrals multiplied to the copula in the system of equations with u . In general, $\delta_k^{[12]} \neq \delta_k^{[21]}$ for $k > 1$ unless in the case of *permutation symmetric* (**isCOP.permsym**) copulas. By theory, $\delta_1^{[12]} = \delta_1^{[21]} = \rho_{\mathbf{C}}/6$ where $\rho_{\mathbf{C}}$ is the *Spearman Rho* **rhoCOP**.

The integral for $\delta_{4;\mathbf{C}}^{[12]}$ does not appear in Brahimi *et al.* (2015) but this and the other forms are verified in the **Examples** and discussion in **Note**. The four $k \in (1, 2, 3, 4)$ for U *wrt* V and V *wrt* U comprise a full spectrum of system of seven (not eight) equations. One equation is lost because $\delta_1^{[12]} = \delta_1^{[21]}$.

Chine and Benatia (2017) describe *trimmed L-comoments* as the multivariate extensions of the univariate *trimmed L-moments* (Elamir and Seheult, 2003) that are implemented in **lmomco**. These are not yet implemented in **copBasic**.

Usage

```
lcomCOP(cop=NULL, para=NULL, as.bilmoms=FALSE, orders=2:5, lo=0, hi=1,
        subdivisions=100L, rel.tol=.Machine$double.eps^0.25, abs.tol=rel.tol,
        stop.on.error=TRUE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
as.bilmoms	A logical to trigger return of the δ_k and the return vectors will be named differently;
orders	The orders of the L-comoments to return, which is internally adjusted if the argument as.bilmoms is set. There is no first order L-comoment and the first index on returned values is set to NA to remain index consistent with the lmomco package. An order greater than 5 is not supported;
lo	Lower limit of integration (must absurdly leave as zero);
hi	Upper integrand (must assuredly leave as unity);
subdivisions	Maximum number of subintervals for <code>integrate()</code> ;
rel.tol	Relative accuracy requested for <code>integrate()</code> ;
abs.tol	Absolute accuracy requested for <code>integrate()</code> ;
stop.on.error	Logical of same name for <code>integrate()</code> to silence for some cases of singularity or other more obscure problems the error “Error in <code>integrate(function(v) : the integral is probably divergent;</code> ” and
...	Additional arguments to pass to the densityCOP function and to <code>integrate()</code> .

Value

An R list of the L-comoments or bivariate L-moments is returned depending on as.bilmoms setting.

bilmomUV	The bivariate L-moments $\delta_k^{[12]}$ of U with respect to V for $k \in [1, 2, 3, 4]$ if orders is 2:5 and there is no NA index as for the L-comoments;
bilmomVU	The bivariate L-moments $\delta_k^{[21]}$ of V with respect to U for $k \in [1, 2, 3, 4]$ if orders is 2:5 and there is no NA index as for the L-comoments;
lcomUV	The L-comoments $\tau_k^{[12]}$ of V with respect to U for $k \in [2, 3, 4, 5]$ if orders is 2:5 and index 1 is NA; and
lcomVU	The L-comoments $\tau_k^{[21]}$ of V with respect to U for $k \in [2, 3, 4, 5]$ if orders is 2:5 and index 1 is NA.

Note

The documentation here is highly parallel to [bilmoms](#) for which that function was developed some years before `lmomCOP` was developed in January 2019. Also, [bilmoms](#) is based on gridded or Monte Carlo integration, and [bilmoms](#) is to be considered **deprecated**. However, it is deliberate that related background and various algorithm testing are still made available in [bilmoms](#). *L-comoment ratio diagrams* are demonstrated by extensive example for many copula in [LCOMDIA_ManyCops](#), [LCOMDIA_GH2cop](#), [LCOMDIA_GH3cop](#), and [ORDSUMcop](#).

Author(s)

W.H. Asquith

References

- Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.
- Brahimi, B., Chebana, F., and Necir, A., 2015, Copula representation of bivariate L-moments—A new estimation method for multiparameter two-dimensional copula models: *Statistics*, v. 49, no. 3, pp. 497–521.
- Chine, Amel, and Benatia, Fatah, 2017, Bivariate copulas parameters estimation using the trimmed L-moments methods: *Afrika Statistika*, v. 12, no. 1, pp. 1185–1197.
- Elamir, E.A.H, and Seheult, A.H., 2003, Trimmed L-moments: *Computational Statistics and Data Analysis*, v. 43, p. 299–314.
- Nelsen, R.B., 2006, *An introduction to copulas*: New York, Springer, 269 p.
- Serfling, R., and Xiao, P., 2007, A contribution to multivariate L-moments—L-comoment matrices: *Journal of Multivariate Analysis*, v. 98, pp. 1765–1781.

See Also

[bilmoms](#), [lcomCOPpv](#), [uvlmoms](#)

Examples

```
## Not run:
para <- list(alpha=0.5, beta=0.93, para1=4.5, cop1=GLcop, cop2=PSP)
lcomCOP(cop=composite2COP, para=para)$lcomUV[3]
# Lcomom:T3[12]= +0.156
lcomCOP(cop=composite2COP, para=para)$lcomVU[3]
# Lcomom:T3[21]= -0.0668
bilmoms(cop=composite2COP, n=10000, para=para, sobol=TRUE)$bilcomoms$T3
# Tau3[12]=+0.1566, Tau3[21]=-0.0655
# The numerical default Monte Carlo integration of bilmoms()
# matches the numerical integration of lcomCOP albeit with a
# substantially slower and less elegant means in bilmoms().
## End(Not run)

## Not run:
# The following Spearman Rho and L-coskew values are predicted for a monitoring
# location of the relation between peak streamflow (V) and time into the
# water year (U) where U and V are "U-statistics."
site_srho <- 0.15536430
site_T3_12 <- 0.03866056
site_T3_21 <- -0.03090144

# Create an objective function for 3D optimization with some explicit intention that
# transforms are used to keep the parameters in acceptable parameter space for the
# alpha [0,1] and beta [0,1] and theta for the Plackett copula and the composite1COP
# used to add two more parameters to the Plackett.
ofunc <- function(par, srho=NA, T3_12=NA, T3_21=NA) {
  alpha <- pnorm(par[1]) # takes -Inf to +Inf ---> 0 to 1 # compositing domain
  beta <- pnorm(par[2]) # takes -Inf to +Inf ---> 0 to 1 # compositing domain
  theta <- exp(par[3]) # takes -Inf to +Inf ---> 0 to +Inf # Plackett domain
```

```

lmr <- lcomCOP(cop=composite1COP,
               para=list(alpha=alpha, beta=beta, para1=theta, cop1=PLcop))
return((lmr$lcomUV[2] - srho )^2 + # look carefully, the 2, 3, 3 index
       (lmr$lcomUV[3] - T3_12)^2 + # use on the lmr list are correct, so do not
       (lmr$lcomVU[3] - T3_21)^2) # expect to see 1, 2, 3 or 2, 3, 4.
}
# initial parameter guess ('middle' [0.5] compositing and independence [1]) and
# showing the transformations involved.
para_init <- c(qnorm(0.5), qnorm(0.5), log(1))
rt <- optim(par=para_init, ofunc,
            srho=site_srho, T3_12=site_T3_12, T3_21=site_T3_21)
lcom_para <- list(alpha=pnorm(rt$par[1]), beta=pnorm(rt$par[2]),
                  para1=exp(rt$par[3]), cop1=PLcop)
sUV <- simCOP(10000, cop=composite1COP, para=lcom_para, col=grey(0, 0.2), pch=16)
# Now as an exercise, consider increasing site_srho or negating it. Consider
# switching the signs on the L-coskews or increasing their magnitudes and study
# the resulting simulation to develop a personal feeling for L-coskew meaning. #
## End(Not run)

```

lcomCOPpv

Simulating the Sample Distribution(s) of L-correlation, L-coskew, and L-cokurtosis for a Copula

Description

EXPERIMENTAL: The function provides two themes of sampling distribution characterization by simulation of the first three L-comoment ratios (L-correlation $\tau_{2[\dots]}$, L-coskew $\tau_{3[\dots]}$ and L-cokurtosis $\tau_{4[\dots]}$) of a copula. Subsequently, the sampling distribution can be used for inference.

First, semi-optional Monte Carlo integration estimation of the L-comoments of the parent copula are computed. Second, simulations involving the sample size n presumed the size of the actual sample from which the estimates of the sample L-comoments given as arguments. These simulations result in a report of the L-moments (not L-comoments) of the sampling distribution and these then are used to compute p-values for the L-comoment matrices provided by the user as a function argument.

Usage

```
lcomCOPpv(n, lcom, cop=NULL, para=NULL, repcoe=5E3, type="gno",
          mcn=1E4, mcrep=10, usemcmu=FALSE, digits=5, ...)
```

Arguments

<code>n</code>	The sample size n . This argument is semi-optional because $n = 0$ can be given to skip corresponding simulations and the <code>nTable</code> on return will only contain NA; this feature permits rapid extraction of the <code>Ntable</code> and thus the <code>lcom</code> contents are simply not used;
<code>lcom</code>	The sample L-comoments (see below);
<code>cop</code>	A copula function;

para	Vector of parameters, if needed, to pass to the copula;
repcoe	The replication coefficient ϕ affecting the number of simulations of size n ;
type	The distribution type used for modeling the distribution of the sampling values. The generalized normal (see distribution type "gno" in package lmomco) accommodates some skewness compared to the symmetry of the normal ("nor") just in case situations arise in which non-ignorable skewness in the sample distribution exists. The distribution abbreviations of package lmomco are recognized for the type argument, but in reality the "nor" and "gno" should be more than sufficient;
mcn	The sample size N passed to the bilmoms function for the Monte Carlo integration. If $N = 0$ then the Monte Carlo integration is not used, otherwise the minimum sample size is internally reset to $N = 4$ so that first four L-moments are computable;
mcrep	The number of replications of the Monte Carlo simulation by bilmoms ;
usemcmu	A logical toggling whether the mean value computed from the replicated Monte Carlo integrations is used instead of the mean values for the small sample simulation for the p-value computations;
digits	The number of digits to round numerical entries in the returned tables and can be NA for no rounding; and
...	Additional arguments to pass to the bilmoms function or to the copula.

Details

The notation $r[\dots]$ refers to two specific types of L-comoment definitions and a blend between the two. The notation $r[12]$ means that the r th L-comoment for random variables $\{X^{(1)}, X^{(2)}\}$ where $X^{(2)}$ is the sorted variable and $X^{(1)}$ is shuffled by the sorting index. Conversely, the notation $r[21]$ means that the r th L-comoment for random variables $\{X^{(1)}, X^{(2)}\}$ where $X^{(1)}$ is the sorted variable and $X^{(2)}$ is shuffled by the sorting index. The notation $r[12 : 21]$ means that the average between the $r[12]$ and $r[21]$ is computed, which might prove useful in circumstances of known or expected symmetry of the L-comoments.

Continuing, $\hat{\tau}_{2[12]}$ is the sample L-correlation, $\hat{\tau}_{3[12]}$ is the sample L-coskew, and $\hat{\tau}_{4[12]}$ is the sample L-cokurtosis all with respect to the sorting of the second variable. The computation of these L-comoment matrices can be made by functions such as function **lcomoms2()** in the **lmomco** package. The number of replications for the simulations involving the n sample size is computed by

$$m = \phi / \sqrt{n},$$

where ϕ is the repcoe replication factor or coefficient. If usemcmu is TRUE then $mcn > 0$ else usemcmu is reset to FALSE.

Value

An R list is returned.

text	A string functioning as a label for the remaining tables;
Ntable	Another R list holding tables of the L-moments of the L-comoments derived from Monte Carlo integration for samples of size $N = mcn$. The simulations are replicated mcrep times; and

ntable Another R list holding tables of the L-moments of the L-comoments derived from the small sample simulations for samples of size $n = n$ as well as the p-values estimated by a generalized normal distribution (see **lmomco** package documentation) of the L-moments using either the small sample means or the mean of the replicated Monte Carlo integrations as dictated by usemcmu. In all circumstances, however, the results for the small sample simulations are tabulated in ntable only the p-value will be reflective the setting of usemcmu.

Note

A significance column for the p-values is added to the right side of the returned ntable and is used to guide the eye in interpretation of results. The significant codes having the following definitions for a two-tailed form:

```
"_" > 0.1;  ".", 0.1;  "*", 0.05;  "**", 0.01;  "***", 0.001
```

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[lcomCOP](#), [COP](#), [kullCOP](#), [vuongCOP](#)

Examples

```
# See Note section of vuongCOP() for an extended discussion of copula inference
## Not run:
Tau <- 0.6410811; para <- GHcop(tau=Tau)$para # This Tau is from a situation of
# two river tributaries. These three L-comoments with univariate L-moments on the
T2 <- c(1, 0.79908960, 0.79908960, 1) # diagonals are derived from those river
# tributaries and downstream of the junction data.
T3 <- c(0, -0.04999318, 0.07689082, 0)
T4 <- c(0, 0.01773833, 0.04756257, 0) # Is the Ho:GHcop rejectable?
LCOM <- list(T2=matrix(T2, nrow=2), T3=matrix(T3, nrow=2), T4=matrix(T4, nrow=2))
set.seed(30312)
ZZ1 <- lcomCOPpv(75, LCOM, cop=GHcop, para=para, repcoe=2000, usemcmu=FALSE)
print(ZZ1)
set.seed(30312)
ZZ2 <- lcomCOPpv(75, LCOM, cop=GHcop, para=para, repcoe=2000, usemcmu=TRUE)
print(ZZ2)
# The results here suggest that the GHcop is not rejectable.
## End(Not run)
```

LCOMDIA_GH2cop

*L-comoment Ratio Diagrams for 2-Parameter Gumbel–Hougaard Extreme Value Copula***Description**

L-comoment ratio diagrams by computation example are shown for the 2-parameter *Gumbel–Hougaard copula* ([GHcop](#)). Diagram computations for the 3-parameter version are found under [LCOMDIA_GH3cop](#). Diagram computations for the 1-parameter version are found among many copulas in [LCOMDIA_ManyCops](#).

Author(s)

W.H. Asquith

See Also[lcomCOP](#), [LCOMDIA_GH3cop](#), [LCOMDIA_ManyCops](#)**Examples**

```
## Not run:
# b1 <- 1 # fixing 2pGHcop gives it L-coskews and L-cokurtoses matching the Clayton.
GH  <- lcomCOP(cop=GHcop, para=c(1, 1.23107734))
clfunc <- function(k, rho=NA) rho - rhoCOP(cop=CLcop, para=exp(k) - 1)
cl  <- exp( uniroot(clfunc, c(-9, +5), rho=GH$lcomUV[2])$root) - 1
CL  <- lcomCOP(cop=CLcop, para=cl) #
## End(Not run)

## Not run:
# -----
# L-comoments of the GH2. Alert, there is a commented out write_feather below, so for
# a "production" run, uncomment that. The reasoning for kicking-out output every nsim
# so that another R session can already be "harvesting" output should the user seek
# to study the output earlier than total run will make.
PLOT <- FALSE
for(f in 1:20) {
  if(PLOT) {
    plot(c(0, 1), c(-0.2, +0.2), type="n", xaxs="i", yaxs="i", las=1,
         xlab="Spearman Rho", ylab="L-coskew (cT3) or L-cokurtosis (cT4)")
    legend("topright", c("L-coskew of GHcop(para=c(b1,b2))",
                        "L-cokurtosis of GHcop(para=c(b1,b2))"), bty="n",
          pch=c(15, 16), pt.cex=0.5, col=c("seagreen4", "orchid3"))
  }
  fileGH2 <- tempfile(); message("Temp File = ", fileGH2)
  nsim <- 10000
  for(i in seq_len(nsim)) {
    if(length(grep("000$", as.character(i)))) message(i,"-", appendLF=FALSE)
    b1 <- 10^(runif(1, min=-5, max=3))+1
    b2 <- ifelse(length(grep("[13579]$", as.character(i))), # 1/2 between 0..1
```

```

      10^runif(1, min=0, max=3), 10^runif(1, min=-6, max=0)) # and 1/2 > 1
GH2 <- lcomCOP(cop=GHcop, para=c(b1, b2), stop.on.error=FALSE)
GH2$lcomUV <- round(GH2$lcomUV, digits=12)
GH2$lcomVU <- round(GH2$lcomVU, digits=12)
d2 <- data.frame(b1, b2,
  T2.12=GH2$lcomUV[2], T3.12=GH2$lcomUV[3], T4.12=GH2$lcomUV[4],
  T2.21=GH2$lcomVU[2], T3.21=GH2$lcomVU[3], T4.21=GH2$lcomVU[4])
row.names(d2) <- NULL; #print(d2)
if(abs(d2$T2.12) < 1 & d2$T2.12 > 0) {
  if(PLOT) points(d2$T2.12, d2$T3.12, cex=0.5, pch=15, col="seagreen4")
  if(PLOT) points(d2$T2.12, d2$T4.12, cex=0.5, pch=16, col="orchid3" )
}
if(abs(d2$T2.21) < 1 & d2$T2.21 > 0) {
  if(PLOT) points(d2$T2.21, d2$T3.21, cex=0.5, pch=15, col="seagreen4")
  if(PLOT) points(d2$T2.21, d2$T4.21, cex=0.5, pch=16, col="orchid3" )
}
k <- i == 1 # triggering mechanism for write.table()
write.table(d2, file=fileGH2, sep="\t", append=! k, col.names=k, row.names=FALSE)
}
message("done")
fGH2 <- read.table(fileGH2, sep="\t", header=TRUE)
file <- paste0("LCM_GH2_", f, ".feather")
message("writing file '", file, "' to ", getwd())
# feather::write_feather(fGH2, file)
}
# -----
files <- list.files(pattern=".feather")
files <- files[grepl("GH2_", files)]
GH2 <- NULL
for(file in files) GH2 <- rbind(GH2, as.data.frame(feather::read_feather(file)))

dr <- 0.001; rhos <- seq(0,1, by=dr); T2 <- c(GH2$T2.12, GH2$T2.21)
T3 <- c(GH2$T3.12, GH2$T3.21); T4 <- c(GH2$T4.12, GH2$T4.21)
T2 <- c(0,0, T2, 1,1); T3 <- c(0,0, T3, 0,0); T4 <- c(0,0, T4, 0,0)
df <- data.frame(T2, T3, T4)
df <- df[df$T2 >= 0,] # giant simulations show no situations of negatives
df$T2[df$T2 > 1] <- 1 # giant simulations show only a few meet this condition and are
# greater than one at the fifth or beyond decimal, so declare as round-off issues and
# not failure itself in the 2-parameter GHcop. (Like 3 in 200,000 simulations)
df <- df[! (df$T2 > 0.997 & df$T3 > -0.468319*df$T2 + 0.468319), ]

plot(df$T2, df$T3, pch=16, cex=0.2, col=grey(0.8), las=1,
  xlab="Spearman Rho", ylab="L-coskew (cT3)")
negT3 <- postT3 <- NULL
for(rho in rhos) {
  negT3 <- c(negT3, min(df$T3[abs(df$T2 - rho) <= dr]))
  postT3 <- c(postT3, max(df$T3[abs(df$T2 - rho) <= dr]))
}
X <- rhos[is.finite(negT3)]; Y <- negT3[is.finite(negT3)]
negT3 <- lm(Y~I(X^ 1)+I(X^ 2)+I(X^ 3)+I(X^ 4)+I(X^ 5)+I(X^ 6)+
  I(X^ 7)+I(X^ 8)+I(X^ 9)+I(X^10)+I(X^11)+I(X^12) )
X <- rhos[is.finite(postT3)]; Y <- postT3[is.finite(postT3)]
posT3 <- lm(Y~I(X^ 1)+I(X^ 2)+I(X^ 3)+I(X^ 4)+I(X^ 5)+I(X^ 6)+

```

```

      I(X^ 7)+I(X^ 8)+I(X^ 9)+I(X^10)+I(X^11)+I(X^12) )
lines(rhos, predict(negT3, newdata=data.frame(X=rhos)), lwd=2)
lines(rhos, predict(posT3, newdata=data.frame(X=rhos)), lwd=2)

t2t3hull <- grDevices::chull(df$T2, df$T3)
polygon(df$T2[t2t3hull], df$T3[t2t3hull], col=NULL, border="seagreen4", lwd=3,lty=2)
write.table(data.frame(T2=df$T2[t2t3hull], T3=df$T3[t2t3hull]),
            file="lcom_GH2cop_nodes.txt", sep="\t")
# The user might consider plotting this polygon() underneath L-comoment curves
# computed and drawn through the recipe in LCOMDIA_ManyCops.Rd.

X <- df$T2[t2t3hull]; Y <- df$T3[t2t3hull]
X <- c(X[Y < 0], 0, 1); Y <- c(Y[Y < 0], 0, 0)
negT3 <- lm(Y~I(X^ 1)+I(X^ 2)+I(X^ 3)+I(X^ 4)+I(X^ 5)+I(X^ 6)+
            I(X^ 7)+I(X^ 8)+I(X^ 9)+I(X^10)+I(X^11)+I(X^12) )
X <- df$T2[t2t3hull]; Y <- df$T3[t2t3hull]
X <- c(X[Y > 0], 0, 1); Y <- c(Y[Y > 0], 0, 0)
posT3 <- lm(Y~I(X^ 1)+I(X^ 2)+I(X^ 3)+I(X^ 4)+I(X^ 5)+I(X^ 6)+
            I(X^ 7)+I(X^ 8)+I(X^ 9)+I(X^10)+I(X^11)+I(X^12) )
plot(df$T2, df$T3, pch=16, cex=0.2, col="lightgreen", las=1,
     xlab="Spearman Rho", ylab="L-coskew (cT3)")
lines(rhos, predict(negT3, newdata=data.frame(X=rhos)), lwd=2)
lines(rhos, predict(posT3, newdata=data.frame(X=rhos)), lwd=2)
print(negT3, 12); print(posT3, 12)
"T3fT2" <- function(rho, type=c("neg", "pos")) {
  type <- match.arg(type)
  if(type == "neg") {
    cs <- c(1.77467318015e-04,
            -3.65537522263e-01, 1.32139171758e+00, -2.26891136600e+01, 2.14717293829e+02,
            -1.20819018654e+03, 4.36103722683e+03, -1.04460882313e+04, 1.67876630052e+04,
            -1.78950352179e+04, 1.21363232163e+04, -4.74068383919e+03, 8.11989541291e+02)
  } else if(type == "pos") {
    cs <- c(-8.77010579087e-06,
            2.45271423943e-01, -2.67994483238e-01, 1.28094964505e+00, -1.28531448606e+01,
            7.61335214074e+01, -2.87746627019e+02, 7.17887377219e+02, -1.19681860869e+03,
            1.31930086424e+03, -9.23025213380e+02, 3.71244634445e+02, -6.53809357075e+01)
  } else {
    stop("should not be here in logic")
  }
  z <- sapply(rho, function(r) sum(sapply(seq_len(length(cs)),
                                          function(p) cs[p] * r^(p-1))))
  return(z)
}
plot(df$T2, df$T3, pch=16, cex=0.2, col="lightgreen", las=1,
     xlab="Spearman Rho", ylab="L-coskew (cT3)")
lines(rhos, T3fT2(rhos, type='neg')) # a last check that the polynomial approximations
lines(rhos, T3fT2(rhos, type='pos')) # have been ported to subroutine acceptably.
# The user might consider plotting these two polynomials underneath L-comoment curves
# computed and drawn through the recipe in LCOMDIA_ManyCops.Rd.
# -----
## End(Not run)

```

LCOMDIA_GH3cop

*L-comoment Ratio Diagrams for 3-Parameter Gumbel–Hougaard Extreme Value Copula***Description**

L-comoment ratio diagrams by computation example are shown for the 3-parameter *Gumbel–Hougaard copula* ([GHcop](#)) with *Marshall–Olkin copula* bounds ([MOcop](#)). Diagram computations for the 2-parameter version are found under [LCOMDIA_GH2cop](#). Diagram computations for the 1-parameter version are found among many copulas in [LCOMDIA_ManyCops](#).

Author(s)

W.H. Asquith

See Also[lcomCOP](#), [LCOMDIA_GH2cop](#), [LCOMDIA_ManyCops](#)**Examples**

```
## Not run:
# -----
# L-comoments of the GH3. Alert, there is a commented out write_feather below, so for
# a "production" run, uncomment that. The reasoning for kicking-out output every nsim
# so that another R session can already be "harvesting" output should the user seek
# to study the output early. Later additions to these Examples are expected to hone
# the results for polynomial boundary and polygonal domain of the relations.
PLOT <- TRUE
for(f in 1:20) {
  if(PLOT) {
    plot(c(0, 1), c(-0.2, +0.3), type="n", xaxs="i", yaxs="i", las=1,
          xlab="Spearman Rho", ylab="L-coskew (cT3) or L-cokurtosis (cT4)")
    legend("topright", c("L-coskew of GHcop(para=c(b1, p2, p3))",
                        "L-cokurtosis of GHcop(para=c(b1, p2, p3))"), bty="n",
          pch=c(3,4), pt.cex=0.5, col=c("seagreen4", "orchid3"))
  }
  fileGH3 <- tempfile(); message("Temp File = ", fileGH3)
  nsim <- 10000
  for(i in seq_len(nsim)) {
    if(length(grep("000$", as.character(i)))) message(i,"-", appendLF=FALSE)
    b1 <- 10^(runif(1, min=-5, max=3))+1
    p2 <- runif(1, min=0, max=1); p3 <- runif(1, min=0, max=1)
    GH3 <- lcomCOP(cop=GHcop, para=c(b1, p2, p3), stop.on.error=FALSE)
    d3 <- data.frame(b1, p2, p3,
                    T2.12=GH3$lcomUV[2], T3.12=GH3$lcomUV[3], T4.12=GH3$lcomUV[4],
                    T2.21=GH3$lcomVU[2], T3.21=GH3$lcomVU[3], T4.21=GH3$lcomVU[4])
    row.names(d3) <- NULL; #print(d3)
    if(abs(d3$T2.12) < 1 & d3$T2.12 > 0) {
      if(PLOT) points(d3$T2.12, d3$T3.12, cex=0.5, pch=15, col="seagreen4")
    }
  }
}
```

```

        if(PLOT) points(d3$T2.12, d3$T4.12, cex=0.5, pch=16, col="orchid3" )
    }
    if(abs(d3$T2.21) < 1 & d3$T2.21 > 0) {
        if(PLOT) points(d3$T2.21, d3$T3.21, cex=0.5, pch=15, col="seagreen4")
        if(PLOT) points(d3$T2.21, d3$T4.21, cex=0.5, pch=16, col="orchid3" )
    }
    k <- i == 1 # triggering mechanism for write.table()
    write.table(d3, file=fileGH3, sep="\t", append=!k, col.names=k, row.names=FALSE)
}
message("done")
fGH3 <- read.table(fileGH3, sep="\t", header=TRUE)
file <- paste0("LCM_GH3_", f, ".feather")
message("writing file '", file, "' to ", getwd())
# feather::write_feather(fGH3, file)
}
# -----
files <- list.files(pattern=".feather")
files <- files[grep("GH3_", files)]
GH3 <- NULL
for(file in files) GH3 <- rbind(GH3, as.data.frame(feather::read_feather(file)))
# -----
plot(GH3$T2.12, GH3$T3.12, cex=0.2, pch=16, col=grey(0.3), las=1,
      xlim=c(0, 0.10), ylim=c(-0.07, 0.0))
abline(0, -0.33) #
## End(Not run)

```

LCOMDIA_ManyCops

*L-comoment Ratio Diagrams for Many Copulas***Description**

L-comoment ratio diagrams by computation example are shown for many copulas of the **copBasic** package.

Author(s)

W.H. Asquith

See Also

[lcomCOP](#), [LCOMDIA_GH2cop](#), [LCOMDIA_GH3cop](#), [ORDSUMcop](#)

Examples

```

## Not run:
# Demonstration of an L-comoment ratio diagram as function of Spearman Rho for various
# copula where each copula is fit to the Spearman Rho and the L-comoments computed.
# -----
# The copulas here are permutation symmetric, which means that T*.12 and T*.21 are
# identical, but plotting for the T*.21 are shown here to make it easier for the user

```

```

# to insert permutation asymmetry say by blending in the breveCOP() or other
# composition frameworks. L-cokurtosis is negated to have it simply plot away from
# the L-coskew.
# -----
LCMempty <- list(lcomUV=rep(NA, 5), lcomVU=rep(NA, 5)) # for comprehensive show of
# features and handling for situations of a Spearman Rho being incompatible with a
# copula, which is the case for the N4212cop that we have reversed here to put the
# tail dependency for moderate Spearman Rho into the upper tail.
rN4212cop <- function(u,v, para, ...) COP(u,v, cop=N4212cop, para=para, reflect=2)

clfunc <- function(k, rho=NA) rho - rhoCOP(cop=COP,
      para=list(reflect=2, cop=CLcop, para=exp(k)-1))
gefunc <- function(k, rho=NA) rho - rhoCOP(cop=gEvcop, para=pnorm(k))
ghfunc <- function(k, rho=NA) rho - rhoCOP(cop=GHcop, para=exp(k)+1)
glfunc <- function(k, rho=NA) rho - rhoCOP(cop=GLcop, para=exp(k) )
hrfunc <- function(k, rho=NA) rho - rhoCOP(cop=HRcop, para=exp(k) )
jofunc <- function(k, rho=NA) rho - rhoCOP(cop=JOcopB5, para=exp(k)+1)
mofunc <- function(k, rho=NA) rho - rhoCOP(cop=M0cop, para=pnorm(rep(k,2)))
n12func <- function(k, rho=NA) rho - rhoCOP(cop=rN4212cop, para=exp(k)+1)
n20func <- function(k, rho=NA) rho - rhoCOP(cop=rN4220cop, para=exp(k) )
rafunc <- function(k, rho=NA) rho - rhoCOP(cop=RAYcop, para=pnorm(k))
rffunc <- function(k, rho=NA) rho - rhoCOP(cop=COP,
      para=list(reflect=2, cop=RFCop, para=pnorm(k)))
D <- NULL; rs <- c(0.0001, 0.0002, 0.0005, 0.0008, 0.001, 0.0013, 0.002, 0.003,
      0.004, 0.005, 0.006, 0.007, 0.008, 0.009, 0.01, 0.015,
      seq(0.02, 0.99, by=0.005), 0.99,
      0.991, 0.992, 0.993, 0.994, 0.995, 0.996, 0.997, 0.998)
rs <- sort(unique(rs)) # to fix defects
for(r in rs) { message(r, "-", appendLF=FALSE) # Spearman Rhos,
  ga <- 2*sin(pi*r/6)
  GA <- lcomCOP(cop=NORMcop, para=ga)
  cl <- exp( uniroot(clfunc, c(-9,+5), rho=r)$root)-1
  CL <- lcomCOP(cop=COP, para=list(cop=CLcop, para=cl, reflect=2))
  fr <- FRcop(rhotau=r, cortype="rho")$para
  FR <- lcomCOP(cop=FRcop, para=fr)
  ge <- pnorm(uniroot(gefunc, c(-4,+4), rho=r)$root)
  GE <- lcomCOP(cop=gEvcop, para=ge)
  gh <- exp( uniroot(ghfunc, c(-10,+4), rho=r)$root)+1
  GH <- lcomCOP(cop=GHcop, para=gh)
  gl <- exp( uniroot(glfunc, c(-9,+4), rho=r)$root)
  GL <- lcomCOP(cop=GLcop, para=gl)
  hr <- exp( uniroot(hrfunc, c(-9,+4), rho=r)$root)
  HR <- lcomCOP(cop=HRcop, para=hr)
  jo <- exp( uniroot(jofunc, c(-10,+4), rho=r)$root)+1
  JO <- lcomCOP(cop=JOcopB5, para=jo)
  mo <- pnorm( uniroot(mofunc, c(-6,+6), rho=r)$root)
  mo <- rep(mo, 2)
  M0 <- lcomCOP(cop=M0cop, para=mo)
  if(r > 0.4784176) { # rhoCOP(rN4212cop, para=1) = 0.4784176
    n12 <- exp( uniroot(n12func, c(-9,+4), rho=r)$root)+1
    N12 <- lcomCOP(cop=rN4220cop, para=n12)
  } else {
    n12 <- NA; N12 <- LCMempty
  }
}

```

```

}
n20 <- exp( uniroot(n20func, c(-10,+4), rho=r)$root)
N20 <- lcomCOP(cop=rN4220cop, para=n20)
ra <- RAYcop(rho=r)
RA <- lcomCOP(cop=RAYcop, para=ra)
rf <- pnorm( uniroot(rffunc, c(-6,6), rho=r)$root)
RF <- lcomCOP(cop=COP, para=list(cop=RFcop, para=rf, reflect=2))
TF <- lcomCOP(cop=Tcop, para=c(ga,1))
x <- data.frame(rho=r, CLpara=cl, FRpara=fr, GApra=ga, GEpara=ge, GHpara=gh,
  GLpara=gl, HRpara=hr, JOpara=jo, MOpara=mo[1],
  N12para=n12, N20para=n20, RApra=ra, RFpara=rf, TFpara=ga,
  CLT2.12=CL$1comUV[2], CLT2.21=CL$1comVU[2],
  CLT3.12=CL$1comUV[3], CLT3.21=CL$1comVU[3],
  CLT4.12=CL$1comUV[4], CLT4.21=CL$1comVU[4],
  FRT2.12=FR$1comUV[2], FRT2.21=FR$1comVU[2],
  FRT3.12=FR$1comUV[3], FRT3.21=FR$1comVU[3],
  FRT4.12=FR$1comUV[4], FRT4.21=FR$1comVU[4],
  GAT2.12=GA$1comUV[2], GAT2.21=GA$1comVU[2],
  GAT3.12=GA$1comUV[3], GAT3.21=GA$1comVU[3],
  GAT4.12=GA$1comUV[4], GAT4.21=GA$1comVU[4],
  GET2.12=GE$1comUV[2], GET2.21=GE$1comVU[2],
  GET3.12=GE$1comUV[3], GET3.21=GE$1comVU[3],
  GET4.12=GE$1comUV[4], GET4.21=GE$1comVU[4],
  GHT2.12=GH$1comUV[2], GHT2.21=GH$1comVU[2],
  GHT3.12=GH$1comUV[3], GHT3.21=GH$1comVU[3],
  GHT4.12=GH$1comUV[4], GHT4.21=GH$1comVU[4],
  GLT2.12=GL$1comUV[2], GLT2.21=GL$1comVU[2],
  GLT3.12=GL$1comUV[3], GLT3.21=GL$1comVU[3],
  GLT4.12=GL$1comUV[4], GLT4.21=GL$1comVU[4],
  HRT2.12=HR$1comUV[2], HRT2.21=HR$1comVU[2],
  HRT3.12=HR$1comUV[3], HRT3.21=HR$1comVU[3],
  HRT4.12=HR$1comUV[4], HRT4.21=HR$1comVU[4],
  JOT2.12=JO$1comUV[2], JOT2.21=JO$1comVU[2],
  JOT3.12=JO$1comUV[3], JOT3.21=JO$1comVU[3],
  JOT4.12=JO$1comUV[4], JOT4.21=JO$1comVU[4],
  MOT2.12=MO$1comUV[2], MOT2.21=MO$1comVU[2],
  MOT3.12=MO$1comUV[3], MOT3.21=MO$1comVU[3],
  MOT4.12=MO$1comUV[4], MOT4.21=MO$1comVU[4],
  N12T2.12=N12$1comUV[2], N12T2.21=N12$1comVU[2],
  N12T3.12=N12$1comUV[3], N12T3.21=N12$1comVU[3],
  N12T4.12=N12$1comUV[4], N12T4.21=N12$1comVU[4],
  N20T2.12=N20$1comUV[2], N20T2.21=N20$1comVU[2],
  N20T3.12=N20$1comUV[3], N20T3.21=N20$1comVU[3],
  N20T4.12=N20$1comUV[4], N20T4.21=N20$1comVU[4],
  RAT2.12=RA$1comUV[2], RAT2.21=RA$1comVU[2],
  RAT3.12=RA$1comUV[3], RAT3.21=RA$1comVU[3],
  RAT4.12=RA$1comUV[4], RAT4.21=RA$1comVU[4],
  RFT2.12=RF$1comUV[2], RFT2.21=RF$1comVU[2],
  RFT3.12=RF$1comUV[3], RFT3.21=RF$1comVU[3],
  RFT4.12=RF$1comUV[4], RFT4.21=RF$1comVU[4],
  TFT2.12=TF$1comUV[2], TFT2.21=TF$1comVU[2],
  TFT3.12=TF$1comUV[3], TFT3.21=TF$1comVU[3],
  TFT4.12=TF$1comUV[4], TFT4.21=TF$1comVU[4] )

```

```

D <- rbind(D, x)
}
Z <- D[1,]
Z$rho <- 0
Z[,grep("[12]$", names(Z))] <- rep(0, length(grep("[12]$", names(Z))))
Z[,grep("para$", names(Z))] <- floor(      Z[,grep("para$", names(Z))])
D <- rbind(Z, D)
Z <- D[1,]
Z$rho <- 1
Z[,grep("[12]$", names(Z))] <- rep(0, length(grep("[12]$", names(Z))))
Z[,grep("para$", names(Z))] <- rep(Inf, length(grep("para$", names(Z))))
D <- rbind( D, Z)
message("done")
row.names(D) <- NULL
D$MOT3.12[is.na(D$MOT3.12)] <- D$MOT3.21[is.na(D$MOT3.12)]
D$TFT3.12[      is.na(D$TFT3.12)] <- approx(D$rho[! is.na(D$TFT3.12)],
      D$TFT3.12[! is.na(D$TFT3.12)],      xout=D$rho[ is.na(D$TFT3.12)])$y
# write.table(D, file="LCOMDIA_ManyCops.txt", sep=",", row.names=FALSE)
# -----
ix <- c(9, 6, 3, 12, 10, 5, 11, 1, 14, 2, 7, 8, 4, 13)
cols <- hcl.colors(length(ix), palette="Dark3")
cols <- cols[ix]
coll <- cols[unlist( sapply(seq_len(length(ix)), function(i) list(rep(i, 2))) )]
rs <- D$rho
plot(rs, D$GHT3.12, ylim=c(-0.20, 0.20), type="n", las=1,
      xlab="Spearman Rho", ylab="L-coskew (cT3) or -negated L-cokurtosis (cT4)")
lines(rs, D$CLT3.12, col=cols[ 1]); lines(rs, -D$CLT4.12, col=cols[ 1], lty=2)
lines(rs, D$CLT3.21, col=cols[ 1]); lines(rs, -D$CLT4.21, col=cols[ 1], lty=2)
lines(rs, D$FRT3.12, col=cols[ 2]); lines(rs, -D$FRT4.12, col=cols[ 2], lty=4)
lines(rs, D$FRT3.21, col=cols[ 2]); lines(rs, -D$FRT4.21, col=cols[ 2], lty=4)
lines(rs, D$GAT3.12, col=cols[ 3]); lines(rs, -D$GAT4.12, col=cols[ 3], lty=2)
lines(rs, D$GAT3.21, col=cols[ 3]); lines(rs, -D$GAT4.21, col=cols[ 3], lty=2)
lines(rs, D$GET3.12, col=cols[ 4]); lines(rs, -D$GET4.12, col=cols[ 4], lty=4)
lines(rs, D$GET3.21, col=cols[ 4]); lines(rs, -D$GET4.21, col=cols[ 4], lty=4)
lines(rs, D$GHT3.12, col=cols[ 5]); lines(rs, -D$GHT4.12, col=cols[ 5], lty=2)
lines(rs, D$GHT3.21, col=cols[ 5]); lines(rs, -D$GHT4.21, col=cols[ 5], lty=2)
lines(rs, D$GLT3.12, col=cols[ 6]); lines(rs, -D$GLT4.12, col=cols[ 6], lty=4)
lines(rs, D$GLT3.21, col=cols[ 6]); lines(rs, -D$GLT4.21, col=cols[ 6], lty=4)
lines(rs, D$HRT3.12, col=cols[ 7]); lines(rs, -D$HRT4.12, col=cols[ 7], lty=2)
lines(rs, D$HRT3.21, col=cols[ 7]); lines(rs, -D$HRT4.21, col=cols[ 7], lty=2)
lines(rs, D$JOT3.12, col=cols[ 8]); lines(rs, -D$JOT4.12, col=cols[ 8], lty=4)
lines(rs, D$JOT3.21, col=cols[ 8]); lines(rs, -D$JOT4.21, col=cols[ 8], lty=4)
lines(rs, D$MOT3.21, col=cols[ 9]); lines(rs, -D$MOT4.21, col=cols[ 9], lty=2)
lines(rs, D$MOT3.12, col=cols[ 9]); lines(rs, -D$MOT4.12, col=cols[ 9], lty=2)
lines(rs, D$N12T3.12, col=cols[10]); lines(rs, -D$N12T4.12, col=cols[10], lty=4)
lines(rs, D$N12T3.21, col=cols[10]); lines(rs, -D$N12T4.21, col=cols[10], lty=4)
lines(rs, D$N20T3.12, col=cols[11]); lines(rs, -D$N20T4.12, col=cols[11], lty=2)
lines(rs, D$N20T3.21, col=cols[11]); lines(rs, -D$N20T4.21, col=cols[11], lty=2)
lines(rs, D$RAT3.12, col=cols[12]); lines(rs, -D$RAT4.12, col=cols[12], lty=4)
lines(rs, D$RAT3.21, col=cols[12]); lines(rs, -D$RAT4.21, col=cols[12], lty=4)
lines(rs, D$RFT3.12, col=cols[13]); lines(rs, -D$RFT4.12, col=cols[13], lty=2)
lines(rs, D$RFT3.21, col=cols[13]); lines(rs, -D$RFT4.21, col=cols[13], lty=2)
lines(rs, D$TFT3.12, col=cols[14]); lines(rs, -D$TFT4.12, col=cols[14], lty=4)

```

```

lines(rs, D$TFT3.21, col=cols[14]); lines(rs, -D$TFT4.21, col=cols[14], lty=4)
txt <- c("cT3 CLcop(reflect)", "-cT4 CLcop(reflect)",
        "cT3 FRcop (equals zero)", "-cT4 FRcop",
        "cT3 NORMcop (equals zero)", "-cT4 NORMcop",
        "cT3 gEVcop", "-cT4 gEVcop",
        "cT3 GHcop", "-cT4 GHcop",
        "cT3 GLcop", "-cT4 GLcop",
        "cT3 HRcop", "-cT4 HRcop",
        "cT3 J0copB5", "-cT4 J0copB5",
        "cT3 M0cop (a=b)", "-cT4 M0cop (a=b)",
        "cT3 rN4212cop", "-cT4 rN4212cop",
        "cT3 rN4220cop", "-cT4 rN4220cop",
        "cT3 RAYcop", "-cT4 RAYcop",
        "cT3 RFcop", "-cT4 RFcop",
        "cT3 Tcop (equals zero)", "-cT4 Tcop")
legend("bottomleft", txt, bty="n", cex=0.7, ncol=4, lty=c(1,2,1,4), col=coll)
mtext("L-comoment ratio diagram for various copula", line=0.8)
# -----
# Now, simulation, note that in samples T[2+].[12|21] are not perfectly equal even for
# known permutation asymmetry. These could be average or used in isolation. Here, we
# just plot the two separately. So, there are double the data points. It takes a lot
# of samples for single sample to look authoritative.
T2 <- T3 <- T4 <- NULL
for(i in 1:60) {
  uv <- simCOP(2000, cop=GLcop, para=0.95, graphics=FALSE)
  lcm <- lmomco::lcomoms2(uv, nmom=4); srho <- (lcm$T2[1,2] + lcm$T2[2,1]) / 2
  points(rep(srho, 2), c(lcm$T3[1,2], lcm$T3[2,1]), cex=0.6, lwd=0.8, pch=24)
  points(rep(srho, 2), -c(lcm$T4[1,2], lcm$T4[2,1]), cex=0.6, lwd=0.8, pch=25)
  T2 <- c(T2, srho)
  T3 <- c(T3, mean(c(lcm$T3[1,2], lcm$T3[2,1])))
  T4 <- c(T4, mean(c(lcm$T4[1,2], lcm$T4[2,1])))
} # BTW : compare the srho to srho <- cor(UV, method="spearman")
mT2 <- mean(T2); mT3 <- mean(T3); mT4 <- mean(T4)
points(mT2, mT3, col=cols[6], lwd=2, bg=grey(0.9), pch=24, cex=2)
points(mT2, -mT4, col=cols[6], lwd=2, bg=grey(0.9), pch=25, cex=2)
# -----
# L-comoment ratio diagram of L-coskew and L-cokurtosis.
# Again both 12 and 21 (UV and VU) are plotted but for permutation symmetry these
# "with respect to's" have the same L-comoments. The GLcop was used in the simulation;
# so, the lines for it are thicker to see that the general centrality of the sample
# L-comoments provide evidence back to the GLcop copula.
plot(D$GHT3.12, xlim=c(0, 0.20), ylim=c(-0.075, 0.075), type="n", las=1,
     xlab="L-coskew (cT3)", ylab="L-cokurtosis (cT4)")
lines( D$CLT3.12, D$CLT4.12, col=cols[ 1])
lines( D$CLT3.21, D$CLT4.21, col=cols[ 1], lty=2)
lines( D$FRT3.12, D$FRT4.12, col=cols[ 2])
lines( D$FRT3.21, D$FRT4.21, col=cols[ 2], lty=4)
lines( D$GAT3.12, D$GAT4.12, col=cols[ 3])
lines( D$GAT3.21, D$GAT4.21, col=cols[ 3], lty=2)
lines( D$GET3.12, D$GET4.12, col=cols[ 4])
lines( D$GET3.21, D$GET4.21, col=cols[ 4], lty=4)
lines( D$GHT3.12, D$GHT4.12, col=cols[ 5])
lines( D$GHT3.21, D$GHT4.21, col=cols[ 5], lty=2)

```

```

lines( D$GLT3.12, D$GLT4.12, col=cols[ 6],          lwd=2)
lines( D$GLT3.21, D$GLT4.21, col=cols[ 6], lty=4, lwd=2)
lines( D$HRT3.12, D$HRT4.12, col=cols[ 7])
lines( D$HRT3.21, D$HRT4.21, col=cols[ 7], lty=2)
lines( D$JOT3.12, D$JOT4.12, col=cols[ 8])
lines( D$JOT3.21, D$JOT4.21, col=cols[ 8], lty=4)
lines( D$MOT3.12, D$MOT4.12, col=cols[ 9])
lines( D$MOT3.21, D$MOT4.21, col=cols[ 9], lty=2)
lines(D$N12T3.12, D$N12T4.12, col=cols[10])
lines(D$N12T3.21, D$N12T4.21, col=cols[10], lty=2)
lines(D$N20T3.12, D$N20T4.12, col=cols[11])
lines(D$N20T3.21, D$N20T4.21, col=cols[11], lty=2)
lines( D$RAT3.12, D$RAT4.12, col=cols[12])
lines( D$RAT3.21, D$RAT4.21, col=cols[12], lty=4)
lines( D$RFT3.12, D$RFT4.12, col=cols[13])
lines( D$RFT3.21, D$RFT4.21, col=cols[13], lty=2)
lines( D$TFT3.12, D$TFT4.12, col=cols[14])
lines( D$TFT3.21, D$TFT4.21, col=cols[14], lty=4)
points( T3, T4, lwd=0.8, cex=0.8)
points(mT3, mT4, lwd=2, cex=2, pch=21, col=cols[6], bg=grey(0.9))
txt <- c("CLcop(reflect)[UV]", "CLcop(reflect)[VU]",
        "FRcop[UV]",          "FRcop[VU]",
        "NORMcop[UV]",        "NORMcop[VU]",
        "gEVcop[UV]",         "gEVcop[VU]",
        "GHcop[UV]",          "GHcop[VU]",
        "GLcop[UV]",          "GLcop[VU]",
        "HRcop[UV]",          "HRcop[VU]",
        "JOcop[UV]",          "JOcop[VU]",
        "MOcop[UV] (a=b)",    "MOcop[VU] (a=b)",
        "rN4212cop[UV]",     "rN4212cop[VU]",
        "rN4220cop[UV]",     "rN4220cop[VU]",
        "RAcop[UV]",         "RAcop[VU]",
        "RFcop[UV]",         "RFcop[VU]",
        "Tcop[UV]",          "Tcop[VU]")
legend("bottomright", txt, cex=0.7, ncol=4, lty=c(1,2), col=coll)
mtext("L-comoment ratio diagram for various copula", line=0.8)
# -----
## End(Not run)

```

lcomoms2.ABcop2parameter

*Convert L-comoments to Parameters of Alpha-Beta Compositions of
Two One-Parameter Copulas*

Description

EXPERIMENTAL—This function converts the *L-comoments* of a bivariate sample to the four parameters of a composition of two one-parameter copulas. Critical inputs are of course the first three dimensionless *L-comoments*: *L-correlation*, *L-coskew*, and *L-cokurtosis*. The most complex input is the *solutionenvir*, which is an environment containing arbitrarily long, but individual tables,

of L-comoment and parameter pairings. These pairings could be computed from the examples in [simcompositeCOP](#).

The individual tables are prescanned for potentially acceptable solutions and the absolute additive error of both L-comoments for a given order is controlled by the tNeps arguments. The default values seem acceptable. The purpose of the prescanning is to reduce the computation space from perhaps millions of solutions to a few orders of magnitude. The computation of the solution error can be further controlled by X or u with respect to Y or v using the comptNerrXY arguments, but experiments thus far indicate that the defaults are likely the most desired. A solution “matching” the L-correlation is always sought; thus there is no uset2err argument. The arguments uset3err and uset4err provide some level of granular control on addition error minimization; the defaults seek to “match” L-coskew and ignore L-cokurtosis. The setreturn controls which rank of computed solution is returned; users might want to manually inspect a few of the most favorable solutions, which can be done by the setreturn or inspection of the returned object from the lcomoms2.cop2parameter function. The examples are detailed and self-contained to the **copBasic** package; curious users are asked to test these.

Usage

```
lcomoms2.ABcop2parameter(solutionenvir=NULL,
                          T2.12=NULL, T2.21=NULL,
                          T3.12=NULL, T3.21=NULL,
                          T4.12=NULL, T4.21=NULL,
                          t2eps=0.1, t3eps=0.1, t4eps=0.1,
                          compt2erruv=TRUE, compt2errvu=TRUE,
                          compt3erruv=TRUE, compt3errvu=TRUE,
                          compt4erruv=TRUE, compt4errvu=TRUE,
                          uset3err=TRUE, uset4err=FALSE,
                          setreturn=1, maxtokeep=1e5)
```

Arguments

solutionenvir	The environment containing solutions;
T2.12	L-correlation $\tau_2^{[12]}$;
T2.21	L-correlation $\tau_2^{[21]}$;
T3.12	L-coskew $\tau_3^{[12]}$;
T3.21	L-coskew $\tau_3^{[21]}$;
T4.12	L-cokurtosis $\tau_4^{[12]}$;
T4.21	L-cokurtosis $\tau_4^{[21]}$;
t2eps	An error term in which to pick a potential solution as close enough on preliminary processing for $\tau_2^{[1\leftrightarrow 2]}$;
t3eps	An error term in which to pick a potential solution as close enough on preliminary processing for $\tau_3^{[1\leftrightarrow 2]}$;
t4eps	An error term in which to pick a potential solution as close enough on preliminary processing for $\tau_4^{[1\leftrightarrow 2]}$;

compt2erruv	Compute an L-correlation error using the 1 with respect to 2 (or u wrt v);
compt2errvu	Compute an L-correlation error using the 2 with respect to 1 (or v wrt u);
compt3erruv	Compute an L-coskew error using the 1 with respect to 2 (or u wrt v);
compt3errvu	Compute an L-coskew error using the 2 with respect to 1 (or v wrt u);
compt4erruv	Compute an L-cokurtosis error using the 1 with respect to 2 (or u wrt v);
compt4errvu	Compute an L-cokurtosis error using the 2 with respect to 1 (or v wrt u);
uset3err	Use the L-coskew error in the determination of the solution. The L-correlation error is always used;
uset4err	Use the L-cokurtosis error in the determination of the solution. The L-correlation error is always used;
setreturn	Set (index) number of the solution to return. The default of 1 returns the preferred solutions based on the controls for the minimization; and
maxtokeep	The value presets the number of rows in the solution matrix. This matrix is filled with potential solutions as the various subfiles of the solutionenvir are scanned. The matrix is trimmed of NAs and error trapping is in place for too small values of maxtokeep. The default value appears appropriate for the feeding of massively large simulated parameter spaces.

Value

An R data.frame is returned.

Author(s)

W.H. Asquith

References

Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.

Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, Extremes in Nature—An approach using copulas: Springer, 289 p.

See Also

[simCOP](#), [simcompositeCOP](#), [composite2COP](#)

Examples

```
## Not run:
# Build an initial parameter to L-comoment mapping table.
mainpara <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
  para1gen=function() { return(10^runif(1, min=-5, max=0)) },
  para2gen=function() { return(10^runif(1, min=0, max=5)) })

nsim <- 1E4
sample.size.for.estimation <- 1000 # really use vastly larger sample size
PlackettPlackettNP <-
```

```

simcompositeCOP(n=sample.size.for.estimation, nsim=nsim, parent=mainpara)
save(PlackettPlackettNP, file="PlackettPlackettNP.RData", compress="xz")

# Plackett-Plackett composited copula from the copBasic package
# Then create an environment to house the "table."
PlackettPlackett <- new.env()
assign("NeedToCreateForDemo", PlackettPlackettNP, envir=PlackettPlackett)
# Now that the table is assigned into the environment, the parameter
# estimation function can be used. In reality, a much much larger
# solution set is needed, but this effort is experimental.

# Now grab the closest Plackett-Plackett solution having the following six
# arbitrary L-comoments. Then simulate 1000 values and plot them to show
# the underlying bivariate distribution.
PPcop <- lcomoms2.ABcop2parameter(solutionenvir=PlackettPlackett,
                                T2.12=-0.5059, T2.21=-0.5110,
                                T3.12= 0.1500, T3.21= 0.1700,
                                T4.12=-0.0500, T4.21= 0.0329,
                                uset3err=TRUE, uset4err=TRUE)

# A user is encouraged to inspect the contents of PPCop to "assess" the
# solution by a method of L-comoments, we will now proceed with showing the
# copula via a simulation of the fitted version.
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, alpha=PPcop$alpha, beta=PPcop$beta,
             para1=PPcop$Cop1Thetas, para2=PPcop$Cop2Thetas)

D <- simCOP(n=5000, cop=composite2COP, para=para, col=rgb(0,0,0,0.1), pch=16)
# The sample L-comoments of the fitted Plackett-Plackett may be found by
lmomco::lmomoms2(D, nmom=4) # from the lmomco package, and six sample values shown
T2.12 <- -0.5151547; T2.21 <- -0.5139863
T3.12 <- 0.1502336; T3.21 <- 0.1721355
T4.12 <- -0.0326277; T4.21 <- 0.0233979
PPcop <- lcomoms2.ABcop2parameter(solutionenvir=PlackettPlackett,
                                T2.12=T2.12, T2.21=T2.21,
                                T3.12=T3.12, T3.21=T3.21,
                                T4.12=T4.12, T4.21=T4.21, uset4err=TRUE)
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, alpha=PPcop$alpha, beta=PPcop$beta,
             para1=PPcop$Cop1Thetas, para2=PPcop$Cop2Thetas)
D <- simCOP(n=5000, cop=composite2COP, para=para, col=rgb(0,0,0,0.1), pch=16)
level.curvesCOP(cop=composite2COP, para=para, delt=.1, ploton=FALSE)
qua.regressCOP.draw(cop=composite2COP, para=para,
                    ploton=FALSE, f=seq(0.05, 0.95, by=0.05))
qua.regressCOP.draw(cop=composite2COP, para=para, wrtV=TRUE,
                    ploton=FALSE, f=seq(0.05, 0.95, by=0.05), col=c(3,2))
diag <- diagCOP(cop=composite2COP, para=para, ploton=FALSE, lwd=4)

image(gridCOP(cop=composite2COP, para=para), col=terrain.colors(20))
# One can inspect alternative solutions like this
# S <- PPCop$solutions$solutions[,1:16]
# B <- S[abs(S$t2.12res) < 0.02 & abs(S$t2.21res) < 0.02 &
#       abs(S$t3.12res) < 0.02 & abs(S$t3.21res) < 0.02, ]
#print(B)
## End(Not run)

```

lcomoms2.ABKGcop2parameter

*Convert L-comoments to Parameters of Alpha-Beta-Kappa-Gamma
Compositions of Two One-Parameter Copulas*

Description

EXPERIMENTAL—This function converts the *L-comoments* of a bivariate sample to the four parameters of a composition of two one-parameter copulas. Critical inputs are of course the first three dimensionless L-comoments: *L-correlation*, *L-coskew*, and *L-cokurtosis*. The most complex input is the `solutionenvir`, which is an environment containing arbitrarily long, but individual tables, of L-comoment and parameter pairings. These pairings could be computed from the examples in [simcompositeCOP](#).

The individual tables are prescanned for potentially acceptable solutions and the absolute additive error of both L-comoments for a given order is controlled by the `tNeps` arguments. The default values seem acceptable. The purpose of the prescanning is to reduce the computation space from perhaps millions of solutions to a few orders of magnitude. The computation of the solution error can be further controlled by *X* or *u* with respect to *Y* or *v* using the `comptNerrXY` arguments, but experiments thus far indicate that the defaults are likely the most desired. A solution “matching” the L-correlation is always sought; thus there is no `uset2err` argument. The arguments `uset3err` and `uset4err` provide some level of granular control on addition error minimization; the defaults seek to “match” L-coskew and ignore L-cokurtosis. The `setreturn` controls which rank of computed solution is returned; users might want to manually inspect a few of the most favorable solutions, which can be done by the `setreturn` or inspection of the returned object from the `lcomoms2.ABKGcop2parameter` function. The examples are detailed and self-contained to the **copBasic** package; curious users are asked to test these.

Usage

```
lcomoms2.ABKGcop2parameter(solutionenvir=NULL,
                             T2.12=NULL, T2.21=NULL,
                             T3.12=NULL, T3.21=NULL,
                             T4.12=NULL, T4.21=NULL,
                             t2eps=0.1, t3eps=0.1, t4eps=0.1,
                             compt2erruv=TRUE, compt2errvu=TRUE,
                             compt3erruv=TRUE, compt3errvu=TRUE,
                             compt4erruv=TRUE, compt4errvu=TRUE,
                             uset3err=TRUE, uset4err=FALSE,
                             setreturn=1, maxtokeep=1e5)
```

Arguments

<code>solutionenvir</code>	The environment containing solutions;
<code>T2.12</code>	L-correlation $\tau_2^{[12]}$;
<code>T2.21</code>	L-correlation $\tau_2^{[21]}$;

T3.12	L-coskew $\tau_3^{[12]}$;
T3.21	L-coskew $\tau_3^{[21]}$;
T4.12	L-cokurtosis $\tau_4^{[12]}$;
T4.21	L-cokurtosis $\tau_4^{[21]}$;
t2eps	An error term in which to pick a potential solution as close enough on preliminary processing for $\tau_2^{[1\leftrightarrow 2]}$;
t3eps	An error term in which to pick a potential solution as close enough on preliminary processing for $\tau_3^{[1\leftrightarrow 2]}$;
t4eps	An error term in which to pick a potential solution as close enough on preliminary processing for $\tau_4^{[1\leftrightarrow 2]}$;
compt2erruv	Compute an L-correlation error using the 1 with respect to 2 (or u wrt v);
compt2errvu	Compute an L-correlation error using the 2 with respect to 1 (or v wrt u);
compt3erruv	Compute an L-coskew error using the 1 with respect to 2 (or u wrt v);
compt3errvu	Compute an L-coskew error using the 2 with respect to 1 (or v wrt u);
compt4erruv	Compute an L-cokurtosis error using the 1 with respect to 2 (or u wrt v);
compt4errvu	Compute an L-cokurtosis error using the 2 with respect to 1 (or v wrt u);
uset3err	Use the L-coskew error in the determination of the solution. The L-correlation error is always used;
uset4err	Use the L-cokurtosis error in the determination of the solution. The L-correlation error is always used;
setreturn	Set (index) number of the solution to return. The default of 1 returns the preferred solutions based on the controls for the minimization; and
maxtokeep	The value presets the number of rows in the solution matrix. This matrix is filled with potential solutions as the various subfiles of the solutionenvir are scanned. The matrix is trimmed of NAs and error trapping is in place for too small values of maxtokeep. The default value appears appropriate for the feeding of massively large simulated parameter spaces.

Value

An R data.frame is returned.

Author(s)

W.H. Asquith

References

- Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.
- Salvadori, G., De Michele, C., Kotegoda, N.T., and Rosso, R., 2007, Extremes in Nature—An approach using copulas: Springer, 289 p.

See Also

[simCOP](#), [simcompositeCOP](#), [composite3COP](#)

Examples

```
## Not run:
mainpara <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
                 para1gen=function() { return(10^runif(1, min=-5, max=5)) },
                 para2gen=function() { return(10^runif(1, min=-5, max=5)) })

nsim <- 1E4
sample.size.for.estimation <- 1000
PlackettPlackettABKGtest <-
  simcomposite3COP(n=sample.size.for.estimation, nsim=nsim, parent=mainpara)
save(PlackettPlackettABKGtest, file="PlackettPlackettABKG.RData", compress="xz")

# Plackett-Plackett composited copula from the copBasic package
# Then create an environment to house the "table".
PlackettPlackettABKG <- new.env()
assign("NeedToCreateForDemo", PlackettPlackettABKGtest, envir=PlackettPlackettABKG)

# Now that the table is assigned into the environment, the parameter estimation
# function can be used. In reality a much much larger solution set is needed.
# Assume one had the following six L-comoments, extract a possible solution.
PPcop <- lcomoms2.ABKGcop2parameter(solutionenvir=PlackettPlackettABKG,
                                   T2.12=-0.5059, T2.21=-0.5110,
                                   T3.12= 0.1500, T3.21= 0.1700,
                                   T4.12=-0.0500, T4.21= 0.0329,
                                   uset3err=TRUE, uset4err=TRUE)

# Now take that solution and setup a parameter object.
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
             alpha=PPcop$alpha, beta=PPcop$beta, kappa=PPcop$kappa, gamma=PPcop$gamma,
             para1=PPcop$Cop1Thetas, para2=PPcop$Cop2Thetas)

# Example Plot Number 1
D <- simCOP(n=2000, cop=composite3COP, para=para, col=rgb(0,0,0,0.1), pch=16)
print(lmomco::lcomoms2(D, nmom=4)) # See the six extracted sample values for this seed.
T2.12 <- -0.4877171; T2.21 <- -0.4907403
T3.12 <- 0.1642508; T3.21 <- 0.1715944
T4.12 <- -0.0560310; T4.21 <- -0.0350028
PPcop <- lcomoms2.ABKGcop2parameter(solutionenvir=PlackettPlackettABKG,
                                   T2.12=T2.12, T2.21=T2.21,
                                   T3.12=T3.12, T3.21=T3.21,
                                   T4.12=T4.12, T4.21=T4.21, uset4err=TRUE)

para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
             alpha=PPcop$alpha, beta=PPcop$beta, kappa=PPcop$kappa, gamma=PPcop$gamma,
             para1=PPcop$Cop1Thetas, para2=PPcop$Cop2Thetas)

# Example Plot Number 2
D <- simCOP(n=1000, cop=composite3COP, para=para, col=rgb(0,0,0,0.1), pch=16)
level.curvesCOP(cop=composite3COP, para=para, delt=0.1, ploton=FALSE)
qua.regressCOP.draw(cop=composite3COP, para=para,
                    ploton=FALSE, f=c(seq(0.05, 0.95, by=0.05)))
```

```

qua.regressCOP.draw(cop=composite3COP, para=para, wrtV=TRUE,
                    ploton=FALSE, f=c(seq(0.05, 0.95, by=0.05)), col=c(3,2))
diag <- diagCOP(cop=composite3COP, para=para, ploton=FALSE, lwd=4)
# Compare plots 1 and 2, some generalized consistency between the two is evident.
# One can inspect alternative solutions like this
# S <- PPCop$solutions$solutions[,1:18]
# B <- S[abs(S$t2.12res) < 0.02 & abs(S$t2.21res) < 0.02 &
#       abs(S$t3.12res) < 0.02 & abs(S$t3.21res) < 0.02, ]
#print(B)
## End(Not run)

```

level.curvesCOP

Compute and Plot Level Curves of a Copula V with respect to U

Description

Compute and plot *level curves* or *level sets* of a copula for V with respect to U (Nelsen, 2006, pp. 12–13). The level curves at levels $t \mapsto [0 + \Delta t, 1 - \Delta t, \Delta t]$ are defined for $U \mapsto [0 + \Delta u, 1 - \Delta u, \Delta u]$ by

$$t \mapsto C(u = U, v),$$

and solving for v . Plotting is provided by this function because level curves are such an important visual attribute of a copula and highly useful for pedagogic purposes. The above equation is implemented by the *inverse of a copula* using [COPinv](#).

Usage

```

level.curvesCOP(cop=NULL, para=NULL, ploton=TRUE, lines=TRUE,
                plotMW=FALSE, ramp=TRUE, delu=0.001, delt=0.10,
                getlevel=NULL, silent=TRUE, ...)

```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
ploton	A logical to toggle on the plot;
lines	A logical to toggle calls to the <code>lines()</code> function in R to draw the lines;
plotMW	A logical to toggle to use the <code>abline()</code> function in R to plot cross lines for the M (M) and W (W) copulas;
ramp	A logical to toggle whether the level curves are ramped in thickness according to the probability of the line. The argument can be used with a <code>lwd</code> setting through the <code>...</code> and can either be a constant, which will override the built-in ramp, or <code>lwd</code> can be defined as a function to replace the built-in ramp in terms of the probability;
delu	The increment for Δu . The default is 1 part in 1,000, which should often provide enough smoothness for many copulas in practice;

delt	The increment Δt for the level curves to plot, defaults to 10-percent intervals. If <code>delt=0.5</code> , then only the median plus the consequences of a defined <code>getlevel</code> is used. If <code>NULL</code> , then a sequence of t values is not made and only <code>getlevel</code> is used (if available);
getlevel	If defined, then it is inserted into the sequence of levels t and that level $t = \text{getlevel}$ is returned in an R list data structure. If more than one level is desired, then instead of repeated calls to this function, the joint.curvesCOP function could be considered;
silent	The argument of the same name given over to <code>try()</code> wrapping the <code>try()</code> operation on forming sequences of t for the curves (see sources); and
...	Additional arguments to pass to the <code>lines()</code> function in R.

Value

Typically no values are returned because this function is used for its side effects, but the arguments can be such that the $\{u, v\}$ for $C(u, v) = t$ are returned within an R list.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[COPinv](#), [level.curvesCOP2](#), [level.setCOP](#), [joint.curvesCOP](#)

Examples

```
## Not run:
level.curvesCOP(cop=M, para=NULL, delt=0.02) # Upper bounds copula
## End(Not run)
## Not run:
D <- level.curvesCOP(cop=P, getlevel=0.56)
str(D) # empty
D <- level.curvesCOP(cop=P, getlevel=0.5)
str(D) # contains stuff
D <- level.curvesCOP(cop=PSP, getlevel=0.8)
str(D) # contains stuff
## End(Not run)
```

level.curvesCOP2

*Compute and Plot Level Curves of a Copula U with respect to V***Description**

Compute and plot *level curves* or *level sets* of a copula for U with respect to V (Nelsen, 2006, pp. 12–13). The level curves at a levels $t \mapsto [0 + \Delta t, 1 - \Delta t, \Delta t]$ are defined for $V \mapsto [0 + \Delta v, 1 - \Delta v, \Delta v]$ by

$$t = C(u, v = V),$$

and solving for u . Plotting is provided by this function because level curves are such an important visual attribute of a copula and highly useful for pedagogic purposes. The above equation is implemented by the *inverse of a copula* using [COPinv2](#).

Usage

```
level.curvesCOP2(cop=NULL, para=NULL, ploton=TRUE, lines=TRUE,
                 plotMW=FALSE, ramp=TRUE, delv=0.001, delt=0.10,
                 getlevel=NULL, silent=TRUE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
ploton	A logical to toggle on the plot;
lines	A logical to toggle calls to the <code>lines()</code> function in R to draw the lines;
plotMW	A logical to toggle to use <code>abline()</code> function in R to plot cross lines for the M (M) and W (W) copulas;
ramp	A logical to toggle whether the level curves are ramped in thickness according to the probability of the line. The argument can be used with a <code>lwd</code> setting through the <code>...</code> and can either be a constant, which will override the built-in ramp, or <code>lwd</code> can be defined as a function to replace the built-in ramp in terms of the probability;
delv	The increment of Δv . The default is 1 part in 1,000, which should often provide enough smoothness for many copulas in practice;
delt	The increment of Δt for the level curves to plot, defaults to 10-percent intervals;
getlevel	If defined and level exists upon stepping through using <code>delt</code> , then the level curve at the <code>getlevel</code> is returned in an R list data structure;
silent	The argument of the same name given over to <code>try()</code> wrapping the <code>try()</code> operation on forming sequences of t for the curves (see sources); and
...	Additional arguments to pass to the <code>lines()</code> function in R.

Value

Typically no values are returned because this function is used for its side effects, but the arguments can be such that the $\{u, v\}$ for $C(u, v) = t$ are returned within an R list.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also[COPinv2](#), [level.curvesCOP](#), [level.setCOP2](#), [joint.curvesCOP2](#)**Examples**

```
## Not run:
level.curvesCOP2(cop=M, para=NULL, delt=0.02) # Upper bounds copula
## End(Not run)
```

level.setCOP

*Compute a Level Set of a Copula V with respect to U***Description**

Compute a *level curve* or *level set* of a copula for V with respect to U (Nelsen, 2006, pp. 12–13). The level curve at level t is defined for $U \mapsto [0 + \Delta u, 1 - \Delta u, \Delta u]$ by

$$t \mapsto \mathbf{C}(u=U, v),$$

and solving for v . The function is largely a dispatcher to features implemented in [level.curvesCOP](#).

Usage

```
level.setCOP(cop=NULL, para=NULL, getlevel=NULL, delu=0.001, lines=FALSE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
getlevel	The level set for t ;
delu	The increment for Δu . The default is 1 part in 1,000, which should often in practice provide enough smoothness for many copulas;
lines	A logical that matches the argument of the same name in level.curvesCOP ; and
...	Additional arguments to pass to the <code>lines()</code> function in <code>R</code> .

Value

The level set for $t = \text{getlevel}$ is returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also[level.setCOP2](#), [level.curvesCOP](#)**Examples**

```
## Not run:
set <- level.setCOP(cop=PSP, getlevel=0.23, delu=0.005)
level.curvesCOP(cop=PSP)
lines(set$U, set$V, col=2, lwd=2) # manually draw the 23rd percentile
set <- level.setCOP(cop=PSP, para=3.1, getlevel=0.67, lines=TRUE, col=4, lwd=4)
# Notice the change in the lines argument and using level.setCOP2 to draw.
mtext("Level Curves and Special Level Sets for PSP copula") #
## End(Not run)
```

level.setCOP2

*Compute a Level Set of a Copula U with respect to V***Description**

Compute a *level curve* or *level set* of a copula for U with respect to V (Nelsen, 2006, pp. 12–13). The level curve at level t is defined for $V \mapsto [0 + \Delta v, 1 - \Delta v, \Delta v]$ by

$$t \mapsto C(u, v=V),$$

and solving for u . The function is largely a dispatcher to features of [level.curvesCOP2](#).

Usage

```
level.setCOP2(cop=NULL, para=NULL, getlevel=NULL, delv=0.001, lines=FALSE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
getlevel	The level set for t ;
delv	The increment for Δv . The default is 1 part in 1,000, which should often in practice provide enough smoothness for many copulas;
lines	A logical that matches the argument of the same name in level.curvesCOP2 ; and
...	Additional arguments to pass to the <code>lines()</code> function in R.

Value

The level set for $t = \text{getlevel}$ is returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[level.setCOP](#), [level.curvesCOP2](#)

Examples

```
## Not run:
set <- level.setCOP2(cop=N4212cop, para=3.1, getlevel=0.23, delu=0.005)
level.curvesCOP2(cop=N4212cop, para=3.1, delv=0.001, delt=0.02)
lines(set$U, set$V, col=2, lwd=2) # manually draw the 23rd percentile
set <- level.setCOP2(cop=N4212cop, para=3.1, getlevel=0.17, lines=TRUE, col=4, lwd=4)
# Notice the change in the lines argument and using levelsetCOP2 to draw.
mtext("Level Curves and Special Level Sets for N4212 copula") #
## End(Not run)
```

LMRuvCOP

Conditional L-moments of the Distribution of U given V

Description

Compute the *conditional L-moments* of U given a V (the Y direction) through the *conditional distribution function* $G(Y)$ using the appropriate *partial derivative* of a copula ($\mathbf{C}(u, v)$) with respect to V . The inversion of the partial derivative is the *conditional quantile function*.

Usage

```
LMRuvCOP(v=seq(0.01, 0.99, by=0.01), cop=NULL, para=NULL, nsim=1E5,
          subdivisions=200L, rel.tol=.Machine$double.eps^0.25, abs.tol=rel.tol, ...)
```

Arguments

<code>v</code>	Nonexceedance probability v in the Y direction;
<code>cop</code>	A copula function with vectorization as in <code>asCOP</code> ;
<code>para</code>	Vector of parameters or other data structures, if needed, to pass to the copula;
<code>nsim</code>	Number of simulations for <i>Monte Carlo integration</i> if numerical integration were to fail;

subdivisions	Argument of same name passed to <code>integrate()</code> ;
rel.tol	Argument of same name passed to <code>integrate()</code> ;
abs.tol	Argument of same name passed to <code>integrate()</code> ; and
...	Additional arguments to pass to derCOP2 .

Value

Value(s) in a list for the median and conditional L-moments given $V = v$ are returned including

median	The conditional median from med.regressCOP2 ;
L1	The conditional mean $\lambda_1 V = v$ from EuvCOP ;
L2	The conditional L-scale $\lambda_2 V = v$;
T3	The conditional L-skew $\tau_3 V = v$;
T4	The conditional L-kurtosis $\tau_4 V = v$;

Author(s)

W.H. Asquith

See Also

[EuvCOP](#), [LMRvuCOP](#), [derCOP2](#)

Examples

```
# Please refer to Examples under LMRvuCOP().
```

LMRvuCOP

Conditional L-moments of the Distribution of V given U

Description

Compute the *conditional L-moments* of V given a U (the X direction) through the *conditional distribution function* $F(X)$ using the appropriate *partial derivative* of a copula ($\mathbf{C}(u, v)$) with respect to U . The conditional L-moments are computed for λ_1 (mean) from [EuvCOP](#) (refer to mathematics therein) and for $r \geq 2$ are computed by

$$\lambda_r = \int_0^1 F(k) \times (1 - F(k)) \times L_r(F(k)) \, dk,$$

where $F(k)$ becomes replaced in **copBasic** with the conditional distribution function of a copula by Nelsen (2006, pp. 13, 40–41) with respect to u :

$$0 \leq \frac{\delta}{\delta u} \mathbf{C}(u, v) \leq 1,$$

which is [derCOP](#) and $L_r(u)$ is defined (Jones, 2004; Asquith, 2011) as

$$L_r(z) = \frac{1}{r-1} \sum_{j=0}^{r-2} (-1)^j \binom{r-1}{j} \binom{r-1}{j+1} z^{r-2-j} (1-z)^j.$$

Asquith (2011) mistakenly lists the leading multiplier $1/(1-r)$ in lieu of $1/(1-r)$ (correct) as shown above; the **lmomco** package contains a suitable errata as discovered in **copBasic** development for LMRvuCOP and [LMRuvCOP](#).

Usage

```
LMRvuCOP(u=seq(0.01, 0.99, by=0.01), cop=NULL, para=NULL, nsim=1E5,
         subdivisions=200L, rel.tol=.Machine$double.eps^0.25, abs.tol=rel.tol, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>cop</code>	A copula function with vectorization as in asCOP ;
<code>para</code>	Vector of parameters or other data structures, if needed, to pass to the copula;
<code>nsim</code>	Number of simulations for <i>Monte Carlo integration</i> if numerical integration were to fail;
<code>subdivisions</code>	Argument of same name passed to <code>integrate()</code> ;
<code>rel.tol</code>	Argument of same name passed to <code>integrate()</code> ;
<code>abs.tol</code>	Argument of same name passed to <code>integrate()</code> ; and
<code>...</code>	Additional arguments to pass to derCOP .

Value

Value(s) in a list for the median and conditional L-moments given $U = u$ are returned including

<code>median</code>	The conditional median from med.regressCOP ;
<code>L1</code>	The conditional mean $\lambda_1 U = u$ from EvuCOP ;
<code>L2</code>	The conditional L-scale $\lambda_2 U = u$;
<code>T3</code>	The conditional L-skew $\tau_3 U = u$;
<code>T4</code>	The conditional L-kurtosis $\tau_4 U = u$;

Author(s)

W.H. Asquith

References

- Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.
- Jones, M.C., 2004, On some expressions for variance, covariance, skewness and L-moments: Journal of Statistical Planning and Inference, v.126, pp. 97–106, doi:[10.1016/j.jspi.2003.09.001](#).
- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[EvuCOP](#), [LMRuvCOP](#), [derCOP](#)

Examples

```
## Not run:
para <- list(cop=breveCOP, para=list(cop=CLcop, para=c(10), breve=0.2, reflect=2))
uv <- simCOP(500, cop=COP, para=para, graphics=FALSE, set.seed=1)

ustar <- 0.96; set.seed(1)
v_special <- replicate(10000, simCOPv(ustar, cop=COP, para=para))
lmr <- lmomco::lmoms(v_special)

u <- v <- seq(0,1, by=0.001)
LMRvu <- LMRvuCOP(u, cop=COP, para=para)
# LMRuv <- LMRuvCOP(v, cop=COP, para=para)
# LMRvu <- LMRuv # swap here to make a quick study of U given V
# uv <- uv[,2:1] # swap here to make a quick study of U given V
# pdf("LMRvuCOP.pdf", height=7, width=7, useDingbats=FALSE)
par(xpd=NA, las=1, lend=1, col.axis="black", cex.axis=0.9,
    mgp=c(2,0.6,0), mai=c(0.65,0.65,0.40,0.65))
plot(uv[,1], uv[,2], type="n", lwd=0.8, col="grey50", bty="n", xlim=c(0,1),
     ylim=c(0,1), xlab="", ylab="", xaxs="i", yaxs="i", xaxt="n", yaxt="n")
lines(rep(ustar, 2), par()$usr[3:4], lty=4, lwd=1.2, col="seagreen4")
par(mgp=c(2,0.3,0), las=0, col.axis="seagreen4", cex.axis=0.8)
txt <- latex2exp::TeX(r'($u^{\star}$')
axis(1, at=ustar, label=txt, lwd=0, lwd.ticks=1.2, tcl=-0.25, col.ticks="seagreen4")
axis(3, at=ustar, label=txt, lwd=0, lwd.ticks=1.2, tcl=-0.25, col.ticks="seagreen4")
axis(1, at=ustar, label=FALSE, lwd=0, lwd.ticks=1.2, tcl=+0.25, col.ticks="seagreen4")
axis(3, at=ustar, label=FALSE, lwd=0, lwd.ticks=1.2, tcl=+0.25, col.ticks="seagreen4")
par(mgp=c(2,0.6,0), las=1, col.axis="black", cex.axis=0.9)
points(uv[,1], uv[,2], lwd=0.9, pch=21, cex=0.8, col="wheat4", bg="wheat1")
axis(1, at=axTicks(1), label=TRUE, lwd=1, lwd.ticks=1, col.ticks="black")
axis(3, at=axTicks(1), label=FALSE, lwd=1, lwd.ticks=1, col.ticks="black")
tix <- seq(0.1, 0.9, by=0.2)
axis(1, at=tix, label=FALSE, lwd=0, lwd.ticks=1, col.ticks="black", tcl=-0.5)
axis(2, at=tix, label=FALSE, lwd=0, lwd.ticks=1, col.ticks="black", tcl=-0.5)
axis(3, at=tix, label=FALSE, lwd=0, lwd.ticks=1, col.ticks="black", tcl=-0.5)
tix <- seq(0.05, 0.95, by=0.05); tix <- tix[grep("5$", as.character(tix))]
axis(1, at=tix, label=FALSE, lwd=0, lwd.ticks=1, col.ticks="black", tcl=-0.2)
axis(2, at=tix, label=FALSE, lwd=0, lwd.ticks=1, col.ticks="black", tcl=-0.2)
axis(3, at=tix, label=FALSE, lwd=0, lwd.ticks=1, col.ticks="black", tcl=-0.2)
mtext("U, nonexceedance probability", side=1, line=1.6)
par(mgp=c(2,0.8,0))
axis(2, at=axTicks(2), label=TRUE, lwd=1, lwd.ticks=1, col.ticks="black")
par(las=0)
mtext("V, nonexceedance probability", side=2, line=2.1)
par(las=1)
lines(u, LMRvu$L1, lwd=2, col="black" ) # mean probability
lines(u, LMRvu$median, lwd=2, col="red1" ) # median probability
lines(u, LMRvu$L2, lwd=2, col="deepskyblue3") # conditional L-scale
points(ustar, mean(v_special), cex=1.7, pch=1, col="black" )
```

```

points(ustar, median(v_special), cex=1.7, pch=1, col="red1" )
points(ustar, lmr$lambda[1], cex=1.3, pch=16, col="black" )
points(ustar, lmr$lambda[2], cex=1.3, pch=16, col="deepskyblue3")
par(new=TRUE)
plot(u, LMRvu$T3, type="n", bty="n", xlim=c(0,1), ylim=c(-1, 1),
      xlab="", ylab="", xaxs="i", yaxs="i", xaxt="n", yaxt="n")
par(col.axis="orchid3", las=0, lend=1); colo <- "orchid3"
mtext("Lskew and Lkurtosis, dimensionless", 4, line=2.15, col=colo)
par(las=1)
axis(4, at=axTicks(2), label=TRUE, lwd=2, lwd.ticks=1, col=colo, col.ticks=colo)
tix <- seq(-0.9, 0.9, by=0.1)
tix <- tix[grepl("[05]$", as.character(tix), invert=TRUE)]
axis(4, at=tix, label=FALSE, lwd=0, lwd.ticks=1, tcl=-0.5, col.ticks=colo)
lines(u, LMRvu$T3, col=colo, lwd=2) # conditional L-skew
lines(u, LMRvu$T4, col=colo, lty=2, lwd=2) # conditional L-kurtosis
points(ustar, lmr$ratios[ 3], cex=1.2, pch=24, col=colo, bg="orchid1")
points(ustar, lmr$ratios[ 4], cex=1.2, pch=25, col=colo, bg="orchid1")
txt <- c(paste0("Conditional (Cond.) mean at ustar = ", ustar, ", EvuCOP(...)",
  "Cond. median at ustar, med.regressCOP(...)",
  "Cond. Lscale (variation) at ustar, LMRvuCOP(...)",
  "Cond. Lskew at ustar, LMRvuCOP(...)",
  "Cond. Lkurtosis at ustar, LMRvuCOP(...)",
  "Demonstration of n=500 sim. values for the copula",
  "Cond. simulated (sim.) (n=10,000) sample mean, mean(...)",
  "Cond. sim. sample median, median(...)",
  "Cond. sim. sample mean, lmoms(...)$lambda[1]",
  "Cond. sim. sample Lscale, lmoms(...)$lambda[2]",
  "Cond. sim. sample Lskew, lmoms(...)$ratios[3]",
  "Cond. sim. sample Lkurtosis, lmoms(...)$ratios[4]"),
  legend("topleft", txt, bty="n", cex=0.6, seg.len=4, y.intersp=1.3,
    lwd=c(2,2,2,2,2, NA, rep(NA, 6)), lty=c(1,1,1,1,2,0,rep(0, 6)),
    pch=c(rep(NA, 5), 21, 1, 1, 16, 16, 24, 25),
    pt.lwd=c(rep(NA,5), 0.9, rep(1,6)),
    pt.cex=c(rep(NA,5), 0.8, 1.7, 1.7, 1.3, 1.3, 1.2, 1.2),
    pt.bg=c(rep(NA,5), "wheat1", rep(NA,4), "orchid1", "orchid1"),
    col=c("black", "red1", "deepskyblue3", colo, colo,
    "wheat4", "black", "red1", "black", "deepskyblue3", colo, colo))
par(xpd=NA, las=1, lend=1, col.axis="black", cex.axis=0.9,
    mgp=c(2,0.6,0), mai=c(0.65,0.65,0.40,0.65))
# dev.off() #
## End(Not run)

```

LzCOPpermsym

Maximum Asymmetry Measure (or Vector) of a Copula by Exchangeability (Permutation Symmetry)

Description

Compute a measure of maximum exchangeable asymmetry of a copula $\mathbf{C}_\Theta$ using *Exchangeability (permutation symmetry)* according to De Baets and De Meyer (2017) by

$$\mu_{\infty \mathbf{C}}^{\text{permsym}} = \mu_{\infty}^{\text{permsym}} = 3 \times \max(|\mathbf{C}_\Theta(u, v) - \mathbf{C}_\Theta(v, u)|)$$

for $(u, v) \in \mathcal{I}^2$. De Baets and De Meyer (2017) comment that among many asymmetric metrics with copulas that $\mu_{\infty}^{\text{permsym}}$ is “by far the most interesting” (De Baets and De Meyer, 2017, p. 36). The 3 multiplier in the definition ensures that $\mu_{\infty}^{\text{permsym}} \in [0, 1]$. De Baets and De Meyer also conclude that permutation symmetry of random variables, in general, is not a desired property in statistical models, and those authors use the μ_{∞} notation in lieu of $L_{\infty}^{\text{permsym}}$ (see documentation related to [LpCOPpermsym](#) within [hoefCOP](#) of *Hoeffding Phi*). The term “Permutation-Mu” is used for this measure in [copBasic-package](#) and in similar contexts.

Usage

```
LzCOPpermsym(cop=NULL, para=NULL, n=5E4,
              type=c("halton", "sobol", "torus", "runif"),
              as.abs=TRUE, as.vec=FALSE, as.mat=FALSE, plot=FALSE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters, if and as needed, to pass to the copula;
n	The simulation size. The default seems sufficient for many practical applications but is suboptimal because the maximum operator in the definition is expected to potentially underestimate the true maximum. When a vector is returned, the default simulation size appears sufficient for many parameter estimation schemes;
type	The type of random number generator on $\mathcal{I}^2$ for computing the maximum (apparent) (see argument n) or a vector of signed differences (see Details);
as.abs	A logical controlling whether the absolute value operation in the $\mu_{\infty}^{\text{permsym}}$ definition is used. This feature permits flexibility retaining the sign of asymmetry;
as.vec	A logical to disable the maximum operation but instead return the a vector of signed differences in the exchanged variables. If this argument is set true, then as.abs will be set false. The return of a vector of signed differences (still multiplied by 3) could be useful in parameter estimation schemes with a similar vector from an <i>empirical copula</i> (EMPIRcop) (see Details);
as.mat	A logical to disable the maximum operation (like as.vec), but instead return a matrix of the $\mathcal{I}^2$ values with third column as the vector of signed differences. If this argument is set true, then as.abs will be set false;
plot	A logical to create a plot of the $\mathcal{I}^2$ domain used in the simulation with a plot title showing the type argument setting;
...	Additional arguments to pass to support flexible implementation.

Details

EFFECT OF RANDOM NUMBER GENERATION—Package **randtoolbox** provides for random number generation on forms different than simply simulating *uniform independent random variables* for $\mathcal{I}^2$. The *Halton*, *Sobol*, and *Torus* types are implemented. The plot argument is useful for the user to see the differences in how these generators canvas the $\mathcal{I}^2$ domain.

The default is Halton, which visually appears to better canvas $\mathcal{I}^2$ without the clumping that simple uniform random variables does and without the larger gaps of Sobol or Torus. Testing indicates that

Halton might generally require the smallest simulation size of the others with simple uniform random variables potentially being the worst and hence such is not the default. Exceptions surely exist depending on the style of the asymmetry. Nevertheless, Halton, Sobol, and Torus produce more consistent estimation behavior with each having a monotone approach towards the true maximum than simple uniform random variables.

The following example is a useful illustration of an asymmetrical *Clayton copula* ($\mathbf{CL}(u, v; \Theta)$, [CLcop](#)) by *composition of a single copula* ([composite1COP](#)) with the theoretical $\mu_{\infty}^{\text{permsym}}$ maxima computed by large sample simulation. A user might explore the effect of the random number generation by changing the type variable.

```
type <- "halton"
para <- list(cop=CLcop, para=20, alpha=0.3, beta=0.1) # asymmetrical Clayton
ti <- LzCOPpermsym(cop=composite1COP, para=para, n=2E6, type=type) # large
ns <- as.integer( 10^seq(1, 4, by=0.05) ) # sequence of simulation sizes
mi <- sapply(ns, function(n) { # produce vector of maxima for simulation size
  LzCOPpermsym(cop=composite1COP, para=para, n=n, type=type) })
ylim <- range(c(0.06, mi, ti)) # vertical limits to ensure visibility
plot(ns, mi, log="x", pch=21, bg=grey(0.9), ylim=ylim, main=type,
      xlab="Simulation size", ylab="Maximum asymmetry measure")
abline(h=ti, lwd=3, col="seagreen") # large sample size estimate in green
```

RELATION TO ANOTHER DISTANCES—The documentation for [LpCOPpermsym](#) lists a supremum definition $L_{\infty}^{\text{permsym}}$, which is like $\mu_{\infty}^{\text{permsym}}$ but lacks the multiplier of 3. However, that documentation mentions a ratio of 1/3 being as upper bounds and hence the De Baets and De Meyer (2017) reasoning for the 3 multiplier to rescale $\mu_{\infty}^{\text{permsym}} \in (0, 1)$. The simple interrelations between the two functions are explored in the following example:

```
para <- c(30, 0.2, 0.95); n <- 5E4
p <- 1
mean(abs(LzCOPpermsym(cop=GHCop, para=para, n=n,
                      as.vec=TRUE)/3)^p)^(1/p) # 0.01867929
      LpCOPpermsym(cop=GHCop, para=para, p=p) # 0.01867940
p <- 3
mean(abs(LzCOPpermsym(cop=GHCop, para=para, n=n,
                      as.vec=TRUE)/3)^p)^(1/p) # 0.02649376
      LpCOPpermsym(cop=GHCop, para=para, p=p) # 0.02649317
```

The critical note is that [LpCOPpermsym](#) is the integral of the absolute differences in permuted differences across $\mathcal{I}^2$. Hence, it is an expectation. The [LzCOPpermsym](#) is difference because of the maximum of the differences. The computations in the example above show how the same information can be extracted from the two functions. De Baets and De Meyer (2017) do not make reference to raising and then rooting by the power p as shown. The examples here provide credence to the default setting of n (simulation size) for several significant figures of similarity.

COPULA PARAMETER ESTIMATION—Parameter estimation using *signed permutation asymmetry vector* can readily be accomplished, which means that we are interested in near-preservation of what permutation asymmetry might be present within the sample. In the self-contained example below, we will assume a parent of *Gumbel–Hougaard* ($\mathbf{GH}(u, v; \Theta)$, [GHcop](#)) extended to asymmetry by using three parameters ($\Theta = (10, 0.8, 0.6)$). Imagine that we unfortunately have a very small

sample size ($n = 100$) as “hundred years of data.” The small sample size facilitates the use of the *checkerboard empirical copula* (**EMPIRCop**); the sample size is small enough that the checkerboard helps smooth through ties. The simulation size for LzCOPpermsym is set “large” as presumed by the existing default. The premise here is that we desire to have near-precise matching to the sample Spearman Rho in conjunction with the permutation asymmetry.

```
# will need packages ** ABCoptim **, ** metaheuristicOpt **, ** pso **
para <- c(10, 0.8, 0.6) # parameters of the parent
nsam <- 100; seed <- 2 # sample size to fit and seed for the RNG
nsim <- 2E4 # note a change from default n argument in LzCOPpermsym()
as.vec <- TRUE # set to FALSE to use just Permutation-Mu
rhoP <- rhoCOP(cop=GHCop, para=para) # parent Spearman Rho
UVsS <- simCOP(cop=GHCop, para=para, n=nsam, seed=seed) # simulate a sample
rhoS <- rhoCOP(as.sample=TRUE, para=UVsS) # sample Spearman Rho
infS <- LzCOPpermsym(cop=EMPIRCop, para=UVsS, n=nsim, type="halton",
  as.vec=as.vec, ctype="checkerboard")
# empirical copula used and returning signed asymmetry vector
# transformation and re-transformation, GHcop paras >1; [0,1]; and [0,1]
tparf <- function(par) c(exp(par[1]) + 1, pnorm( par[2] ), pnorm( par[3] ))
rparf <- function(par) c(log(par[1] - 1), qnorm( par[2] ), qnorm( par[3] ))

ofunc <- function(par, norho=FALSE, usetrans=FALSE) { # objective function
  if(usetrans) { mypara <- tparf(par) } else { mypara <- par }
  rhoT <- rhoCOP(cop=GHCop, para=mypara) # simulated Spearman Rho
  infT <- LzCOPpermsym(cop=GHCop, para=mypara, n=nsim, type="halton",
    as.vec=as.vec)
  err <- mean( (infT - infS)^2 ) # mean squared errors
  ifelse(norho, err, (rhoT - rhoS)^2 + err) # with Spearman Rho or not
} # norho and usetrans are inaccessible for metaOpt() call to come later
# ofunc defaults are chosen to keep us from writing two objective functions

init.par <- c(2, 0.5, 0.5) # initial parameters in real space
tnit.par <- rparf(init.par); rt <- NULL # initial parameter guess
try( rt <- optim(tnit.par, ofunc, norho=FALSE, usetrans=TRUE) )
# 3D optimization with unbounded parameters using transformation.
if(is.null(rt)) stop("fatal, optim() returned NULL")
# construct GHcop parameters from optimization with re-transformation
sara <- tparf(rt$par)
rhoT <- rhoCOP(cop=GHCop, para=sara) # theoretical Spearman Rho
UVsT <- simCOP(cop=GHCop, para=sara, n=nsam, seed=seed, # same seed sim by
  cex=0.3, pch=16, col="red", ploton=FALSE) # est. parameters
# maximum likelihood
mara <- mleCOP(UVsS, cop=GHCop, init.para=tnit.par, parafn=tparf)$para

# metaheuristic methods follow and need lower and upper parameter bounds
lb <- c(1, 0, 0); ub <- c(100, 1, 1) # lower and upper parameter bounds
# Artificial Bee Colony optimization
# https://en.wikipedia.org/wiki/Artificial_bee_colony_algorithm
be <- ABCoptim::abc_optim(init.par, ofunc, norho=FALSE, lb=lb, ub=ub)
```

```

sara_bee <- be$par
# Particle Swarm optimization
# https://en.wikipedia.org/wiki/Particle_swarm_optimization
ps <- pso::psoptim(init.par, ofunc, norho=FALSE, lower=lb, upper=ub)
sara_pso <- ps$par

# Genetic Algorithm optimization
# https://en.wikipedia.org/wiki/Genetic_algorithm
rngv <- matrix(c(lb, ub), nrow=2, byrow=TRUE)
ctrl <- list(numPopulation=30) # fairly slow running, so 30 for speed
mo <- metaheuristicOpt::metaOpt(ofunc, algorithm="GA",
                                numVar=3, rangeVar=rngv, control=ctrl)
sara_alt <- mo$result

level.curvesCOP(cop=GHcop, para=para)
level.curvesCOP(cop=GHcop, para=mara,   ploton=FALSE, col="blue"   )
level.curvesCOP(cop=GHcop, para=sara,   ploton=FALSE, col="red"     )
level.curvesCOP(cop=GHcop, para=sara_bee, ploton=FALSE, col="orchid3" )
level.curvesCOP(cop=GHcop, para=sara_pso, ploton=FALSE, col="seagreen")
level.curvesCOP(cop=GHcop, para=sara_alt, ploton=FALSE, col="wheat"  )

```

Comparison of level curves between the known parent, the parameter estimation using function LzCOPpermsym, and the *maximum likelihood* by [mleCOP](#) shows that signed asymmetry differences can be used for parameter estimation. One could use the maximum as in the definition, but for purposes of high-dimensional optimization, using the vector might be better to prevent local minima (less optimal solutions) being found if the $\mu_{\infty}^{\text{permsym}}$ was used. Because vectors of differences between empirical copula and the fitted copula are involved, measures of fit using such differences are expected to be more favorable to optimization than using LzCOPpermsym than perhaps maximum likelihood. Some measures of fit like RMSE ([rmseCOP](#)), for example, are often, smaller for the sara fitted parameters than for the mara fitted (maximum likelihood). The maximum likelihood approach should favor towards smaller AIC ([aicCOP](#)) and BIC ([bicCOP](#)). Finally, setting `as.vec <- FALSE`, re-running, and thus using $\mu_{\infty}^{\text{permsym}}$, will likely show parameter estimates, visible through the level curves, that are much less favorable.

```

n <- 1E6; nm <- c("Parent", "MLE", "SARA", "ABC", "PSO", "GA")
AIC <- c(aicCOP(UVsS, cop=GHcop, para=para   ),
        aicCOP(UVsS, cop=GHcop, para=mara   ),
        aicCOP(UVsS, cop=GHcop, para=sara   ),
        aicCOP(UVsS, cop=GHcop, para=sara_bee),
        aicCOP(UVsS, cop=GHcop, para=sara_pso),
        aicCOP(UVsS, cop=GHcop, para=sara_alt))
RHO <- c(rhoCOP(      cop=GHcop, para=para   ),
        rhoCOP(      cop=GHcop, para=mara   ),
        rhoCOP(      cop=GHcop, para=sara   ),
        rhoCOP(      cop=GHcop, para=sara_bee),
        rhoCOP(      cop=GHcop, para=sara_pso),
        rhoCOP(      cop=GHcop, para=sara_alt)); names(RHO) <- nm
LZP <- c(LzCOPpermsym(cop=GHcop, para=para,   n=n, type="halton"),
        LzCOPpermsym(cop=GHcop, para=mara,   n=n, type="halton"),

```

```

LzCOPpermsym(cop=GHcop, para=sara, n=n, type="halton"),
LzCOPpermsym(cop=GHcop, para=sara_bee, n=n, type="halton"),
LzCOPpermsym(cop=GHcop, para=sara_pso, n=n, type="halton"),
LzCOPpermsym(cop=GHcop, para=sara_alt, n=n, type="halton"))
names(AIC) <- nm; names(RHO) <- nm; names(LZP) <- nm
print(AIC)
print(RHO)
print(LZP) # approximate results are shown in the table below
#      Parent      MLE      SARA      ABC      PSO      GA
# AIC  -928.5670 -916.2741 -939.8767 -939.8861 -939.8860 -937.8460
# RHO   0.6132539 0.5433072 0.6058879 0.6058639 0.6058639 0.6058761
# LZP   0.1076471 0.1000950 0.1186881 0.1187269 0.1187269 0.1211495

```

Value

A scalar value for the measure is returned or other as dictated by arguments.

Author(s)

W.H. Asquith

References

De Baets, B., and De Meyer, H., 2017, Chapter 3—A look at copulas in a curved mirror: New York, Springer, ISBN 978–3–319–64221–5, pp. 33–47.

See Also

[LpCOPpermsym](#), [isCOP.permsym](#)

Examples

```

LzCOPpermsym(cop=PSP) # 0, permutation symmetric
LzCOPpermsym(cop=GHcop, para=c(10, 0.9, 0.3)) # 0.17722861

# See also results of
# isCOP.permsym(cop=PSP) # TRUE
# isCOP.permsym(cop=GHcop, para=c(10, 0.9, 0.3)) # FALSE

## Not run:
sapply(1:4, function(r) { # Four rotations of a Galambos copula
  Lz <- LzCOPpermsym(cop=COP, para=list(cop=GLcop, para=2, reflect=r))
  UV <- simCOP(1000, cop=COP, para=list(cop=GLcop, para=2, reflect=r))
  mtext(paste0("Reflection ", r, " : Permutation-Mu =", Lz)); Lz })
# [1] 0.00000000 0.00000000 0.07430326 0.07430326
## End(Not run)

## Not run:
# Example purpose = exploring sample sizes permutation symmetry metrics to
# stabilize by some examples of difference metrics for empirical copula that
# are computed in about the simplest and most efficient manner at the expense

```

```

# of the copying of UV inside the two EMPIRcop() calls, which could become
# substantial if n ---> large, but we do not require additional book keeping
# for u <- c(UV[,1], UV[,2]); v <- c(UV[,2], UV[,1]);
# h <- EMPIRcop(u,v, para=UV) and then h[first half] - h[second half].
DF <- NULL; para <- list(breve=0.2, para=10, cop=GHCop)
for(n in as.integer(10^seq(log10(100), log10(50000), by=0.04))) {
  UV <- simCOP(n, cop=breveCOP, para=para, graphics=FALSE)
  hh <- EMPIRcop(UV[,1], UV[,2], para=UV) - EMPIRcop(UV[,2], UV[,1], para=UV)
  mad <- mean(abs(hh)); rmse <- sqrt(mean(hh^2)); max <- 3*max(hh)
  DF <- rbind(DF, data.frame(n, mad, rmse, max))
}
ylim <- c(0, max(sapply(2:ncol(DF), function(i) max(DF[,i]))))
plot( DF$n, DF$mad, col="red", type="l", log="x", ylim=ylim)
lines(DF$n, DF$rmse, col="seagreen")
lines(DF$n, DF$max, col="blue") #
## End(Not run)

```

M

The Fréchet–Hoeffding Upper-Bound Copula

Description

Compute the *Fréchet–Hoeffding upper-bound copula* (Nelsen, 2006, p. 11), which is defined as

$$\mathbf{M}(u, v) = \min(u, v).$$

This is the copula of perfect association (*comonotonicity*, *perfectly positive dependence*) between U and V and is sometimes referred to as the *comonotonicity copula*. Its opposite is the $\mathbf{W}(u, v)$ copula (*countermonotonicity copula*; [W](#)), and statistical *independence* is the $\mathbf{\Pi}(u, v)$ copula ([P](#)).

Usage

```
M(u, v, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[W](#), [P](#)

Examples

```
M(0.4, 0.6)
M(0, 0)
M(1, 1)
```

med.regressCOP	<i>Perform Median Regression using a Copula by Numerical Derivative Method for V with respect to U</i>
----------------	--

Description

Perform *median regression* (Nelsen, 2006, pp. 217–218) of a copula by inversion of numerical derivatives of the copula ([derCOPinv](#)). The documentation for [qua.regressCOP](#) provides mathematical details. The [qua.regressCOP.draw](#) supports so-called *quantile regression* along various probability levels (see **Examples**).

Usage

```
med.regressCOP(u=seq(0.01, 0.99, by=0.01), cop=NULL, para=NULL, level=NA, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>level</code>	The level of the prediction interval to compute. For example, <code>level=0.95</code> will compute the 95-percent prediction interval as will <code>level=0.05</code> because internally a reflection check is made; and
<code>...</code>	Additional arguments to pass such qua.regressCOP and derCOPinv that are called in succession.

Value

An R `data.frame` of the median regressed probabilities of V and provided U values is returned. Note: if `level` is used, then the column ordering of the returned `data.frame` changes—please access the columns by the named idiom. The lower- and upper-prediction interval is contained in the columns respectively titled `Vlwr` and `Vupr` to mimic nomenclature somewhat of function `predict.lm()` in R.

Note

An extended demonstration is needed concerning prediction intervals by median regression and a comparison to well-known linear regression. This also affords an opportunity to have **copBasic** interact with the **copula** package to gain access to the *Gaussian copula* and a *maximum pseudo-likelihood* estimation of the parameter.

First, a function `NORMcop()` is defined to form the interconnect between the two packages. It is critically important that the user recognize that the so-called copula object as built by the **copula** package is treated as the canonical `para` argument in **copBasic** calls herein.

```
"NORMcop" <- # pCoupla() from package copula is analogous to COP()
function(u,v, para=NULL, ...) {
  if(length(u) == 1) u <- rep(u, length(v)) # see asCOP() for reasoning of
  if(length(v) == 1) v <- rep(v, length(u)) # this "vectorization" hack
  return(copula::pCopula(matrix(c(u,v), ncol=2), para))
}
```

The parameter $\Theta \in [-1, 1]$ (*Pearson R*) and $\rho_C(\Theta)$ (*Spearman Rho*, **rhoCOP**) and $\tau_C(\Theta)$ (*Kendall Tau*, **tauCOP**) are according to Salvadori *et al.* (2007, p. 255) the values

$$\rho_C(\Theta) = \frac{2}{\pi} \arcsin(\Theta)$$

and

$$\tau_C(\Theta) = \frac{6}{\pi} \arcsin(\Theta/2).$$

Second, a bivariate *Gaussian copula* is defined with a parameter $\Theta = 0.7$ (thus $\rho_C = 0.6829105$, `rhoCOP(NORMcop, norm.cop)`) and then $n = 255$ samples simulated from it. These are then cast into standard normal variates to mimic the idea of bivariate data in nonprobability units and facilitation regression comparison.

```
norm.cop <- copula::normalCopula(c(0.7), dim = 2) # define a Gaussian copula
UVs      <- copula::rCopula(255, norm.cop) # draw 255 samples
UVs      <- as.data.frame(UVs); X <- qnorm(UVs[,1]); Y <- qnorm(UVs[,2])
```

Third, the *Weibull plotting positions* from the `pp()` function of package **lmomco** are used to estimate the empirical probabilities of the data in UV that are casted into an **R** matrix because the **copula** package expects the data as a matrix for the default parameter estimation. The code is completed by the specification of the fitted *Gaussian copula* in `fnorm.cop`.

```
UV <- as.matrix(data.frame(U=lmomco::pp(X, sort=FALSE),
                          V=lmomco::pp(Y, sort=FALSE)))
para <- copula::fitCopula(copula::normalCopula(dim=2), UV)
para <- summary(para)$coefficients[1] # maximum pseudo-likelihood est.
fnorm.cop <- copula::normalCopula(para, dim=2)
```

Fourth, ordinary-least-squares (OLS) linear regression for $Y | X$ and $X | Y$ is computed, and the results plotted on top of the data points. The 2/3rd-prediction limits are computed by `predict.lm()` and also shown.

```
# Classical linear regressions from two perspectives.
LMyx <- lm(Y~X);      LMxy <- lm(X~Y)
YonX <- summary(LMyx); XonY <- summary(LMxy)

QUorV <- seq(-3,3, by=0.05) # vector for graphical operations
plot(X, Y, col=8, pch=21)
lines(QUorV, YonX$coefficients[1]+YonX$coefficients[2]*QUorV, col=2, lwd=4)
tmp <- predict.lm(LMyx, list(X=X), level=2/3, interval="prediction")
lines(X, tmp[,2], col=2); lines(X, tmp[,3], col="red" )

lines(XonY$coefficients[1]+XonY$coefficients[2]*QUorV, QUorV, col=3, lwd=4)
tmp <- predict.lm(LMxy, list(Y=Y), level=2/3, interval="prediction")
lines(tmp[,2], Y, col="green"); lines(tmp[,3], Y, col="green")
```

Finally, the demonstration ends with plotting of the median regression for the Gaussian copula and drawing the regression lines. The two median regression lines are nearly coincident with the OLS regression lines as anticipated with a reasonably large sample size albeit maximum pseudo-likelihood was used to estimate the copula parameter. The mean of a uniform distributed variable given say $U = u$ (horizontal axis) is $1/2$, which coincides with the median. The median regression lines thus are coincident with the OLS lines even though OLS and real-space (native units of X and Y) were not used for their computation. The $2/3$ -prediction intervals also are plotted for comparison.

```
UorV <- c(0.001, seq(.02,0.98, by=.02), 0.999)
MEDreg <- med.regressCOP( u=UorV, level=2/3, cop=NORMcop, para=fnorm.cop)
MEDreg2 <- med.regressCOP2(v=UorV, level=2/3, cop=NORMcop, para=fnorm.cop)
lines(qnorm(UorV), qnorm(MEDreg$V), col="blue", lty=2)
lines(qnorm(UorV), qnorm(MEDreg$Vlwr), col="blue", lty=2)
lines(qnorm(UorV), qnorm(MEDreg$Vupr), col="blue", lty=2)
lines(qnorm(MEDreg2$U), qnorm(UorV), col="magenta", lty=2)
lines(qnorm(MEDreg2$Ulw), qnorm(UorV), col="magenta", lty=2)
lines(qnorm(MEDreg2$Uupr), qnorm(UorV), col="magenta", lty=2)
```

A curious aside (Joe, 2014, p. 164) about the *Gaussian copula* is that *Blomqvist Beta* (**blomCOP**) is equal to *Kendall Tau* (**tauCOP**), which can be checked by

```
blomCOP(cop=NORMcop, para=norm.cop) # 0.4936334
tauCOP( cop=NORMcop, para=norm.cop) # 0.493633
```

and obviously the $\beta_C = \tau_C$ are numerically the same.

Author(s)

W.H. Asquith

References

- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kotegoda, N.T., and Rosso, R., 2007, Extremes in Nature—An approach using copulas: Springer, 289 p.

See Also

[med.regressCOP2](#), [qua.regressCOP](#), [qua.regressCOP.draw](#)

Examples

```
## Not run:
# INDEPENDENCE YIELDS STRAIGHT LINES, RED IS THE MEDIAN REGRESSION
FF <- seq(0.1, 0.9, by=0.1)
plot(c(0,1),c(0,1), type="n", lwd=3,
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
# Draw the regression of V on U and then U on V (wrtV=TRUE)
qua.regressCOP.draw(f=FF, cop=P, para=NA, ploton=FALSE)
qua.regressCOP.draw(f=FF, cop=P, para=NA, ploton=FALSE, wrtV=TRUE, lty=2)#
## End(Not run)

## Not run:
# NEGATIVE PLACKETT THETA YIELDS CURVES DOWN TO RIGHT, RED IS THE MEDIAN REGRESSION
theta <- 0.5; FF <- seq(0.1, 0.9, by=0.1)
plot(c(0,1),c(0,1), type="n", lwd=3,
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
# Draw the regression of V on U and then U on V (wrtV=TRUE)
qua.regressCOP.draw(f=FF, cop=PLACKETTcop, ploton=FALSE, para=theta)
qua.regressCOP.draw(f=FF, cop=PLACKETTcop, ploton=FALSE, para=theta, wrtV=TRUE, lty=2)#
## End(Not run)
```

med.regressCOP2

Perform Median Regression using a Copula by Numerical Derivative Method for U with respect to V

Description

Perform *median regression* of a copula (Nelsen, 2006, pp. 217–218) by inversion of numerical derivatives of the copula ([derCOPinv2](#)). The documentation for [qua.regressCOP2](#) provides mathematical details.

Usage

```
med.regressCOP2(v=seq(0.01,0.99, by=0.01), cop=NULL, para=NULL, level=NA, ...)
```

Arguments

<code>v</code>	Nonexceedance probability v in the Y direction;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>level</code>	The level of the prediction interval to compute. For example, <code>level=0.95</code> will compute the 95-percent prediction interval as will <code>level=0.05</code> because internally a reflection check is made; and
<code>...</code>	Additional arguments to pass such qua.regressCOP2 and derCOPinv2 that are called in succession.

Value

An R `data.frame` of the median regressed probabilities of U and provided V values is returned. Note: if `level` is used, the column ordering of the returned `data.frame` changes—please access the columns by the named idiom. The lower- and upper-prediction interval bounds are contained in the columns respectively titled `U1wr` and `Uupr` to mimic nomenclature somewhat of function `predict.lm()` in R.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[med.regressCOP](#), [qua.regressCOP2](#), [qua.regressCOP.draw](#)

Examples

```
# See examples under med.regressCOP
```

mleCOP	<i>Maximum Pseudo-Log-Likelihood Estimation for Copula Parameter Estimation</i>
--------	---

Description

Perform maximum pseudo-log-likelihood estimation (pMLE) for copula parameters by maximizing the function:

$$\mathcal{L}(\Theta_p) = \sum_{i=1}^n \log[c(F_x(x_i), F_y(y_i); \Theta_p)],$$

where $\mathcal{L}(\Theta_p)$ is the log-likelihood for parameter vector Θ_p of dimension p , and $c(u, v; \Theta_p)$ is the bivariate copula density. The u and v are estimated by the respective empirical cumulative distribution functions $u = F_x(\dots)$ and $v = F_y(\dots)$ for each of the joint realizations of a sample of size n . The $c(u, v)$ is numerically estimated by the copula using the [densityCOP](#) function.

Usage

```
mleCOP(u, v=NULL, cop=NULL, parafn=function(k) return(k),
       interval=NULL, init.para=NULL, verbose=FALSE, control=list(),
       the.zero=.Machine$double.eps^0.25, s=0, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction and if <code>NULL</code> then <code>u</code> is treated as a two column <code>R</code> data.frame;
<code>cop</code>	A copula function;
<code>parafn</code>	A function responsible for generating the parameters. This is often just a simple return of a parameter vector as copBasic uses this style of parameterization, but this function can take over parameter remapping to handle boundary conditions to benefit the search or provide an interface into other copula packages in <code>R</code> (see Examples);
<code>interval</code>	The search interval for root finding, by <code>stats::optimise()</code> , if the parameter dimension of the copula is $p = 1$. The interval is not used for $p \geq 2$;
<code>init.para</code>	The initial guesses for the parameters for the p -dimensional optimization for $p \geq 2$. The initial guess is used, by <code>stats::optim()</code> , if the parameter dimension of the copula is $p = 1$ and <code>interval</code> is <code>NULL</code> (see Examples);
<code>verbose</code>	A logical that internally is converted to integer to trigger 1 (sum of logs of densityCOP shown), 2 (add reporting of the copula parameter on each iteration), or more levels of verbose reporting scheme within the objective function. This is independent from the <code>control\$trace</code> of function <code>optim()</code> ;
<code>control</code>	This argument is the argument of the same name for <code>optim()</code> ;
<code>the.zero</code>	The value for “the zero” of the copula density function. This argument is the argument of the same name for densityCOP . The default here is intended to suggest that a tiny nonzero value for density will trap the numerical zero densities;
<code>s</code>	A vector of at least two presumably uniformly distributed or regular sequence of nonexceedance probabilities in U for simulation of V by simCOPv and plotting of these U and V . This plotting is only made if the length of s is nonzero and <code>verbose</code> is greater than or equal to 2. This plotting feature for the s is pedagogical and intended for demonstration or teaching opportunities. This feature has no utility for the optimization itself; and
<code>...</code>	Additional arguments to pass, see source code for the internally used functions that can pick these additional arguments up.

Value

The value(s) for the estimated parameters are returned within an `R` list where the elements listed below are populated unique to this package. The other elements of the returned list are generated from either the `optimise()` ($1D, p = 1$) or `optim()` ($pD, p \geq 2$) functions of `R`.

<code>para</code>	The parameter(s) in a canonical element after the one-dimensional root finding ($p = 1$) or multi-dimensional optimization ($p \geq 2$) solutions are passed through <code>parafn</code> so that these are in the parameter units of the copula and not necessarily those transformed for the optimization;
<code>packagetext</code>	A helpful message unique to the copBasic package;

loglik	The maximum of the log-likelihood matching the name for the same quantity by the function <code>fitCopula</code> in package copula though a separate implementation is used in copBasic ;
AIC	Akaike information criterion (AIC) (see also aicCOP): $AIC = 2p - 2\mathcal{L}(\Theta_p)$; and
BIC	Bayesian information criterion (BIC) (see also bicCOP): $BIC = p \log(n) - 2\mathcal{L}(\Theta_p)$.

Note

This section provides for a more thorough assessment of pMLE than shown in the **Examples**.

INTERFACE TO THE COPULA PACKAGE—A not uncommon question to the author is how can **copBasic** support copulas from other packages? A **copBasic** pMLE implementation to the *Gaussian copula* from the **copula** package is thus useful for instruction.

Two interface functions are required for the pMLE situation. First, interface the **copula** package in a generic form for the **copBasic** package:

```
"cB2copula" <- # pCoupla() from package copula is analogous to COP()
function(u,v, para=NULL, ...) {
  if(length(u) == 1) u <- rep(u, length(v)) # see asCOP() for reasoning of
  if(length(v) == 1) v <- rep(v, length(u)) # this "vectorization" hack
  return(copula::pCopula(matrix(c(u,v), ncol=2), para))
}
```

where the `para` argument above must be built by the features of the **copula** package. The following function then provides for parameter setup specific to the *Gaussian copula* having parameter ρ :

```
copula2cBpara <- function(rho) return(copula::normalCopula(rho, dim = 2))
```

Now, let us perform a parameter estimate for a sample of size $n = 900$:

```
set.seed(162); UV <- simCOP(n=900, cop=cB2copula, para=copula2cBpara(0.45))
mleCOP(UV, cop=cB2copula, parafn=copula2cBpara, interval=c(-1,1))$para
# rho.1 = 0.4248822
```

The search interval for the *Gaussian copula* is $\rho \in [-1, 1]$, and the final result is $\rho = 0.4458822$.

MULTI-DIMENSIONAL EXAMPLE OF pMLE—Consider a 2-parameter *Gumbel–Hougaard copula* (**GH**(Θ_1, Θ_2)) but now use the `parafn` argument to provide boundary condition assistance through function `GH2pfunc` to the `optim()` function that performs the maximization.

```
set.seed(162); UV <- simCOP(n=890, cop=GHcop, para=c(2.4, .06))
GH2pfunc <- function(p) { return(c(exp(p[1])+1, exp(p[2]))) }
ML <- mleCOP(UV$U, UV$V, cop=GHcop, init.para=c(1,1), parafn=GH2pfunc)
print(ML$para) # [1] 2.2755018 0.1194788
```

and the result is $\Theta_{1,2} = (2.2755018, 0.1194788)$. Next, consider now a 3-parameter **GH**(Θ, π_1, π_2) copula and again use the `parafn` argument through function `GH3pfunc` but notice that the 2nd and 3rd parameters are now mapped into $0 \leq \pi_1, \pi_2 \leq 1$ domain using the `pnorm()` function.

```
set.seed(162); UV <- simCOP(n=500, cop=GHcop, para=c(5.5, .6, .9))
GH3pfunc <- function(p) { return(c(exp(p[1])+1, pnorm(p[2]), pnorm(p[3]))) }
ML <- mleCOP(UV$U, UV$V, cop=GHcop, init.para=c(1, .5, .5), parafn=GH3pfunc)
print(ML$para) # [1] 5.3742229 0.6141652 0.9382638
```

and the result is $\Theta = 5.3742229$ and $\pi_{1,2} = (0.6141652, 0.9382638)$.

ANOTHER MULTI-DIMENSIONAL EXAMPLE OF $pMLE$ —Finally, an experiment can be made fitting a 3-parameter $\mathbf{GH}(\Theta, \pi_1, \pi_2)$ to a simulation from a 2-parameter $\mathbf{GH}(\beta_1, \beta_2)$, where the seed is just arbitrary and the *Vuong Procedure* (`vuongCOP`) is used to compare fits and make inference. The parameter functions `GH2pfunc` and `GH3pfunc` are as before.

```
set.seed(10); UV <- simCOP(n=500, cop=GHcop, para=c(1.7, 1.86))
GH2pfunc <- function(p) { return(c(exp(p[1])+1, exp(p[2])) ) }
GH3pfunc <- function(p) { return(c(exp(p[1])+1, pnorm(p[2]), pnorm(p[3])) ) }
para1 <- mleCOP(UV, cop=GHcop, init.para=c(1,1), parafn=GH2pfunc)$para
para2 <- mleCOP(UV, cop=GHcop, init.para=c(1,.5,.5), parafn=GH3pfunc)$para
vuongCOP(UV, cop1=GHcop, para1=para1, cop2=GHcop, para2=para2)$message
#[1] "Copula 1 has better fit than Copula 2 at 100 x (1-alpha) level"
```

The results show the 2-p $\mathbf{GH}$ is a better fit to the simulated data than the 3-p $\mathbf{GH}$, which seems a bit self evident? Plot some same-seeded simulations just to confirm.

```
set.seed(67) # First the estimated parameters but with the correct model.
UV <- simCOP(n=200, GHcop, para=para1, snv=TRUE, pch=16, col=2)
set.seed(67) # Second, the estimated incorrect model.
UV <- simCOP(n=200, GHcop, para=para2, snv=TRUE, ploton=FALSE)
```

Yes, differences in form are manifest in the produced graphic. Now, let us try another set of parameters and again an arbitrarily-chosen seed.

```
set.seed(10); UV <- simCOP(n=500, cop=GHcop, para=c(1.91, 0.16))
para1 <- mleCOP(UV, cop=GHcop, init.para=c(1,1), parafn=GH2pfunc)$para
para2 <- mleCOP(UV, cop=GHcop, init.para=c(1,.5,.5), parafn=GH3pfunc)$para
vuongCOP(UV, cop1=GHcop, para1=para1, cop2=GHcop, para2=para2)$message
#[1] "Copulas 1 and 2 are not significantly different at 100 x (1-alpha)"
```

The results show equivalence, let us now check a graphic.

```
set.seed(67); z <- simCOP(n=200, GHcop, para=para1, snv=TRUE, pch=16, col=2)
set.seed(67); z <- simCOP(n=200, GHcop, para=para2, snv=TRUE, ploton=FALSE)
```

The differences are small but the differences might be inflating into the lower left corner. What sample size could conceivably begin to distinguish between the copula?

```
kullCOP(cop1=GHcop, cop2=GHcop, para1=para1, para2=para2) # 625 on this run

nsim <- 20; set.seed(67)
Results <- sapply(1:nsim, function(i) {
  UV <- simCOP(n=625, cop=GHcop, para=c(1.91, .16), graphics=FALSE)
```

```
p1 <- mleCOP(UV, cop=GHcop, init.para=c(1,1),      parafn=GH2pfunc)$para
p2 <- mleCOP(UV, cop=GHcop, init.para=c(1,.5,.5), parafn=GH3pfunc)$para
vuongCOP(UV, cop1=GHcop, para1=p1, cop2=GHcop, para2=p2)$result })
sum(Results)
```

The summation yields 6 of 20 for which copula 1 has the better fit, but with $n = 1,000$ instead of $n = 625$, the sum of the Results is 13 of 20 (so better than half the time). This seems to be in conflict with what the n_{fg} sample size from [kullCOP](#) should be telling. The author thinks it should be 18 to 19 of 20 (95th percentile) based on what the [kullCOP](#) is reported to do (NEED TO LOOK INTO THIS).

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[densityCOP](#)

Examples

```
# See also extended code listings and discussion in the Note section

## Not run:
# Here, we study the trajectory of the objective function in a simple
# 1-dimensional optimization. See how we must provide the interval.
set.seed(162); UV <- simCOP(n=188, cop=PLcop, para=5.6)
ML <- mleCOP(UV$U, UV$V, cop=PLcop, interval=c(0.1, 40)) # 5.225459 estimated

Thetas <- 10^(seq(log10(0.001), log10(100), by=0.005))
MLs <- sapply(Thetas, function(k)
  densityCOP(UV$U, UV$V, cop=PLcop, para=k, sumlogs=TRUE))
plot(Thetas, MLs, log="x", type="l", # draw the pMLE solution process
     xlab="Plackett Theta", ylab="sum of log densities")
lines(rep(ML$para, 2), c(ML$objective, par()$usr[3]), col="red")
points(ML$para, ML$objective, pch=16, col="red") #
## End(Not run)

## Not run:
# Here, we study again 1-dimensional optimization but use the
# multidimensional version with an alert issued.
set.seed(149); UV <- simCOP(1000, cop=CLcop, para=pi)
# Warning messages about using optim() for 1D solution
mleCOP(UV, cop=CLcop, init.para=2)$para # 3.082031
# No warning message, optimise() called instead.
mleCOP(UV, cop=CLcop, interval=c(0,1E2))$para # 3.081699
## End(Not run)
```

```
## Not run:
# Here, we evaluate a 2-dimensional problem using a Plackett again but with
# the addition of asymmetry towards high V outliers from the Plackett cloud.
# This example also adds the internal verbose and graphic diagnostics for
# the iterations of the optimizer. Here, we learn that we need on a time have
# some idea where the solution might lay so that we can provide a suitable
# set of initial parameters for the algorithm.
para <- list(beta=-0.1, cop=PLcop, para1=1000)
UV <- simCOP(2000, cop=breveCOP, para=para); abline(0, 1, col="red", lwd=3)
PL2pfunc <- function(p) { # see here example of parameter transform
  list(beta=2*pnorm(p[1])-1, para=exp(p[2]), cop=PLcop) # [-1,+1], >=0
}
init.para <- c(0.2535, log(0.02)) # These will not find a solution with this
# guess of negative association, but the next works by using an external
# estimate of the Plackett parameters and here we test with a positive
# skewness (beta for breveCOP > 0) although we know the parent had negative.
init.para <- c(0.2535, log(PLACKETTpar(UV$U, UV$V, byrho=TRUE))) # beta=0.200
rt <- mleCOP(u=UV$U, v=UV$V, init.para=init.para, cop=breveCOP,
  parafn=PL2pfunc, verbose=2, s=seq(0,1, by=0.005)) #
## End(Not run)
```

MOcop

The Marshall–Olkin Copula

Description

The *Marshall–Olkin copula* (Dobrowolski and Kumar, 2014, eq. 2.6; Nelsen, 2006, p. 53), which is also known as the *Generalized Cuadras–Augé copula*, having parameters $\alpha, \beta \in (0, 1)$ is given by

$$C_{(\alpha, \beta)}(u, v) = \min(u^{1-\alpha}v, uv^{1-\beta}),$$

and efficiently computed by compound equations for $u^\alpha \geq v^\beta$

$$C_{(\alpha, \beta)}(u, v) = \mathbf{MO}(u, v) = u^{1-\alpha}v,$$

and for $u^\alpha < v^\beta$ is

$$C_{(\alpha, \beta)}(u, v) = \mathbf{MO}(u, v) = uv^{1-\beta}.$$

As $\alpha, \beta \rightarrow 1$, the copula limits to the *comonotonicity coupla* ($\mathbf{M}(u, v)$; **M**), and as $\alpha, \beta \rightarrow 0$, the copula limits to the *independence copula* ($\mathbf{II}(u, v)$; **P**). The copula can be expected to have a visible singularity as the parameters increase and (or) a simulation size becomes large. The copula with $\alpha = \beta$ is *permutation symmetric* (**isCOP.permsym**, **breveCOP**) and is known as the *Cuadras–Augé copula*. *Spearman Rho* (**rhoCOP**) of the copula is

$$\rho_C = 3\alpha\beta / (2\alpha + 2\beta - \alpha\beta),$$

and Kendall tau ([tauCOP](#)) is

$$\tau_C = \alpha\beta/(\alpha + \beta - \alpha\beta).$$

Parameter estimation using ρ_C and τ_C is problematic because the two statistics are so “correlated” with each other and that either nonunique solutions or nonconverged solutions manifest with simultaneous numerical optimization. Also, application of *maximum likelihood* ([mleCOP](#)) can be problematic because of varying contributions of singularities that the copula can produce. The total contribution of singularity is given by $S(u \rightarrow 1, v \rightarrow 1) = \alpha\beta/(\alpha + \beta - \alpha\beta)$, which Nelsen (2006, p. 165) remarks that it is “interesting to note that” Kendall tau is the singular component. Lastly, the *L-comoments* of copulas ([lcomCOP](#)) appear an attractive mechanism for parameter estimation not effected by singularities (refer to **Examples**). The MO copula has a role in the three-parameter *Gumbel–Hougaard copula* ([GHcop](#)).

Usage

```
MOcop(u, v, para=NULL, lcomCOP=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the α, β parameters of the copula;
<code>lcomCOP</code>	A vector containing ρ_C and the two L-coskews (refer to Examples) from which the parameters are returned; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned. Otherwise if either `lcomCOP` is given, then the α, β are computed by numerical optimization the the Spearman Rho and the two L-coskews (refer to package **lmomco** and function `lmomco::lcomoms2` for more details) (refer the demonstration of L-comoment parameter estimation in the **Examples** below).

Author(s)

W.H. Asquith

References

- Dobrowolski, E., and Kumar, P., 2014, Some properties of the Marshall–Olkin and generalized Cuadras–Augé families of copula: Australian Journal of Mathematical Analysis and Applications: v. 11, no. 1, art. 2, pp. 1–13, accessed on August 10, 2025, at <https://ajmaa.org/searchroot/files/pdf/v11n1/v11i1p2.pdf>.
- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[P](#), [taildepCOP](#)

Examples

```
## Not run:
# **** Demonstration of copula fitting by Method of L-comoments ****
n <- 300 # large enough sample size to clearly show L-comoment viability
UV <- simCOP(n, cop=M0cop, seed=1, para=c(0.3, 0.7), pch=16, col="seagreen")
lcm <- lmomco::lcomoms2(UV) # sample L-comoments (mean off diag. T2 == Spearman Rho)
sara <- M0cop(lcomCOP=c(mean(diag(lcm$T2[2:1,1:2])), lcm$T3[1,2], lcm$T3[2,1]))
sara$para <- round(sara$para, digits=8)
uv <- simCOP(n, cop=M0cop, seed=1, para=sara$para, col="blue", ploton=FALSE)
message("M0cop : alpha=", sara$para[1], ", beta=", sara$para[2])
mara <- mleCOP(UV[,1], v=UV[,2], cop=M0cop, init.para=c(0.2,0.2),
               parafn=function(x) pnorm(x))
mara$para <- round(mara$para, digits=8)
uv <- simCOP(n, cop=M0cop, seed=1, para=mara$para, col="red", ploton=FALSE)
message("M0cop : alpha=", mara$para[1], ", beta=", mara$para[2]) #
## End(Not run)

## Not run:
# This example developed as a demonstrator of the disarray that the
# Spearman Footrule measures as a nonunique relation to Spearman Rho.
# The permutation asymmetry causes the relation between the two to
# "paint" for positive association as shown in the graphic produced.
cex <- c(1.3, 1.0, 0.7, 0.4)
col <- c("#7D0112", "#67C7B1", "#1F28A2", "#CCC86C")
pch <- c(1, 16, 1, 16)
plot(c(-0.5, 1), c(0, 1), type="n", xlab="Spearman Footrule", ylab="")
mtext(paste0("Absolute value of Spearman rho\n",
             "Actual sign is the same as for the Footrule"), side=2, line=2)
mtext(paste0("Attainable Marshall-Olkin copula, M0cop() with permutation\n",
             "inclusion (breveCOP) and reflections (COP)"), line=1)
for(r in 1:4) { # reflections, COP
  for(k in seq(-1, 1, by=0.1)) { # permutation asymmetry insertion, breveCOP
    for(a in seq(0, 1, by=0.11)) { # increment slightly different from beta
      for(b in seq(0, 1, by=0.09)) { # increment slightly different from alpha
        para <- list(cop=M0cop, para=c(a,b), breve=k, reflect=r)
        fotT <- footCOP(cop=breveCOP, para=para)
        rhoT <- rhoCOP(cop=breveCOP, para=para)
        points(fotT, abs(rhoT), lwd=0.8, pch=pch[r], cex=cex[r], col=col[r])
        #UV <- simCOP(100, cop=breveCOP, para=para)
      }
    }
  }
} #
## End(Not run)
```

Description

Compute shuffles of *Fréchet–Hoeffding upper-bound copula* (Nelsen, 2006, p. 173), which is defined by partitioning $\mathbf{M}$ within $\mathcal{I}^2$ into n subintervals:

$$\mathbf{M}_n(u, v) = \min\left(u - \frac{k-1}{n}, v - \frac{n-k}{n}\right)$$

for points within the partitions

$$(u, v) \in \left[\frac{k-1}{n}, \frac{k}{n}\right] \times \left[\frac{n-k}{n}, \frac{n-k+1}{n}\right], k = 1, 2, \dots, n$$

and for points otherwise outside the partitions

$$\mathbf{M}_n(u, v) = \max(u + v - 1, 0).$$

The support of $\mathbf{M}_n$ consists of n line segments connecting coordinate pairs $\{(k-1)/n, (n-k)/n\}$ and $\{k/n, (n-k+1)/n\}$ as stated by Nelsen (2006). It is useful that Nelsen stated such as this helps to identify that Nelsen's typesetting of the terms in the second square brackets—the V direction—is reversed from that shown in this documentation. The *Spearman Rho* ([rhoCOP](#)) is defined by $\rho_C = (2/n^2) - 1$, and the *Kendall Tau* ([tauCOP](#)) by $\tau_C = (2/n) - 1$.

Usage

```
M_N5p12b(u, v, para=1, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A positive integer $n \in 1, 2, \dots$; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[M](#), [ORDSUMcop](#), [W_N5p12a](#)

Examples

```

M_N5p12b(0.4, 0.6, para=3)

## Not run:
# Nelsen (2006, exer. 5.12b, p. 173, fig. 5.3b)
UV <- simCOP(1000, cop=M_N5p12b, para=4) #
## End(Not run)

```

N4212cop

*The Copula of Equation 4.2.12 of Nelsen's Book***Description**

The *N4212 copula* (Nelsen, 2006, p. 91; eq. 4.2.12) is named by the author (Asquith) for the **copBasic** package and is defined as

$$C_{N4212}(u, v; \Theta) = \left(1 + [(u^{-1} - 1)^{\Theta} + (v^{-1} - 1)^{\Theta}]^{1/\Theta} \right)^{-1}.$$

The **N4212**(u, v) copula is *not comprehensive* because for $\Theta = 1$ the copula becomes the so-called **PSP**(u, v) copula (see [PSP](#)) and as $\Theta \rightarrow \infty$ the copula becomes **M**(u, v) (see [M](#)). The copula is undefined for $\Theta < 1$. The N4212 copula has respective *lower-* and *upper-tail dependency* (see [taildepCOP](#)).

Although **copBasic** is intended to not implement or “store house” the enormous suite of copula functions available in the literature, the N4212 copula is included to give the package another copula to test or test in numerical examples.

Usage

```
N4212cop(u, v, para=NULL, infis=100, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula;
<code>infis</code>	What is infinity? Testing shows that <code>infis = $\Theta > 100$</code> is about right to consider the copula as becoming M (u, v) (see M); and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

Examples

```
N4212cop(0.4,0.6, para=1) == PSP(0.4,0.6) # TRUE
N4212cop(0.4,0.6, para=10) # 0.3999928
taildepCOP(cop=N4212cop, para=10) # LamL = 0.93303; LamU = 0.92823
## Not run:
D <- simCOP(n=400, cop=N4212cop, para=2)
D <- simCOP(n=400, cop=N4212cop, para=10, ploton=FALSE, col=2)
D <- simCOP(n=400, cop=N4212cop, para=100, ploton=FALSE, col=3)#
## End(Not run)
```

N4220cop

The Copula of Equation 4.2.20 of Nelsen's Book

Description

The *N4220 copula* (Nelsen, 2006, p. 118; eq. 4.2.20) is named by the author (Asquith) for the **copBasic** package and is defined as

$$C_{N4220}(u, v; \Theta) = \left[\log \left(\exp(u^{-\Theta}) + \exp(v^{-\Theta}) - \exp(1) \right) \right]^{-1/\Theta}.$$

The **N4220**(u, v) copula is *not comprehensive* because for $\Theta = 0^+$ the copula becomes the *independence* **P**(u, v) copula (**P**) and as $\Theta \rightarrow \infty$ the copula becomes **M**(u, v) (**M**). The copula is undefined for $\Theta < 0$. This copula has *lower-tail dependency*, which is the reverse of the **rN4220**(u, v) that has *uppr-tail dependency*. Hence, a reflected (refer to **COP**, reflect=2) version of this copula is natively supported as the **rN4220cop** to put the tail dependency into the upper-right corner and more “hydrologic-system like.”

Usage

```
N4220cop(u, v, para=NULL, insertM=TRUE, ...)
```

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction;
para	A vector (single element) of parameters—the Θ parameter of the copula;
insertM	For the tail that pinches out towards M (u, v) (M) use an <i>ad hoc</i> algorithm to insert that copula into the tail (refer to source code) and that algorithm is dependent on the simCOP and the numerical partial derivative inversion for simulation that the package is based around; and
...	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

Examples

```
sapply(c(0.1, 1, 10), function(p) N4220cop(0.4, 0.6, para=p))
# 0.2611771 0.3714941 0.4000000 # perfect correlation controls for p=10
```

NORMcop	<i>The Normal (Gaussian) Copula</i>
---------	-------------------------------------

Description

The *Normal copula (Gaussian copula)* (Salvadori *et al.*, 2007, pp. 255–256) is

$$C_{\Theta}(u, v) = \text{NORM}(u, v; \Theta) = \int_{-\infty}^{\Phi^{(-1)}(u)} \int_{-\infty}^{\Phi^{(-1)}(v)} \frac{1}{2\pi\sqrt{1-\Theta^2}} \exp\left(-\frac{s^2 - 2\Theta st + t^2}{2(1-\Theta^2)}\right) ds dt,$$

where $\Theta \in [-1, 1]$ and $\Phi^{(-1)}(x)$ is the quantile function of the *univariate normal distribution*. The copula, as $\Theta \rightarrow -1^+$, limits to the *countermonotonicity copula* (**W**(u, v); **W**), as $\Theta \rightarrow 0$ limits, to the *independence coupla* (**P**(u, v); **P**), and as $\Theta \rightarrow 1^-$, limits to the *comonotonicity copula* (**M**(u, v); **M**). The copula has *lower-tail* and *upper-tail dependency parameters* equal to zero, but such is not true for the closely related *t-Student copula* (**Tcop**). The *Spearman Rho* (**rhoCOP**) is $\rho_C = (6/\pi) \cdot \text{asin}(\Theta/2)$ and *Kendall Tau* (**tauCOP**) is $\tau_C = (2/\pi) \cdot \text{asin}(\Theta)$. The parameter Θ is readily computed by $\Theta = 2 \cdot \sin(\pi \cdot \rho_C/6)$ or by $\Theta = \sin(\pi \cdot \tau_C/2)$.

Usage

```
NORMcop(u, v, para=NULL, rho=NULL, tau=NULL, fit=c("rho", "tau"), ...)
```

Arguments

- u Nonexceedance probability u in the X direction;
- v Nonexceedance probability v in the Y direction;
- para A vector (single element) of parameters—the Θ parameter of the copula;
- rho Optional Spearman Rho from which the parameter will be estimated and presence of rho trumps tau;

<code>tau</code>	Optional Kendall Tau from which the parameter will be estimated;
<code>fit</code>	If <code>para</code> , <code>rho</code> , and <code>tau</code> are all <code>NULL</code> , then the <code>u</code> and <code>v</code> represent the sample. The measure of association by the <code>fit</code> declaration will be computed and the parameter estimated subsequently. The <code>fit</code> has no other utility than to trigger which measure of association is computed internally by the <code>cor</code> function in R ; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned. Otherwise if either `rho` or `tau` is given, then the Θ is computed and a list having

<code>para</code>	The parameter Θ ;
<code>rho</code>	Spearman Rho if the <code>rho</code> is given; and
<code>tau</code>	Kendall Tau if the <code>tau</code> is given but also if both <code>rho</code> and <code>tau</code> are <code>NULL</code> as mentioned next.

and if `para=NULL` and `rho` and `tau=NULL`, then the values within `u` and `v` are used to compute Spearman Rho (`fit="rho"`) or Kendall Tau (`fit="tau"`) and then compute the parameter, and this is returned in the aforementioned list.

Note

A mimic of the wrappers on `mvtnorm::pmvnorm()` from the **copula** package (v1.1-6) are made within **NORMcop**. The rest of the implementation is unique to **copBasic**. An implementation of a function in the **copBasic** style that would natively interconnect to **copula** package version of the copula could be as follows:

```
"NORMcop" <-      # pCoupla() from package copula is analogous to COP()
function(u,v, para=NULL, ...) {
  if(length(u) == 1) u <- rep(u, length(v)) # see asCOP() for reasoning of
  if(length(v) == 1) v <- rep(v, length(u)) # this "vectorization" hack
  para <- copula::normalCopula(              c(para), dim=2)
  return( copula::pCopula(matrix(c(u,v), ncol=2), para) )
}
```

Author(s)

W.H. Asquith

References

Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, *Extremes in Nature—An approach using copulas*: Springer, 289 p.

See Also

[Tcop](#)

Examples

```
## Not run:
para <- NORMcop(para=NULL, rho=0.9, taildep=0.3) # 0.907981
UV <- simCOP(1000, cop=NORMcop, para=para$para) # compare to similar Tcop example
## End(Not run)
```

ORDSUMcop

Ordinal Sums of M-Copula

Description

Compute *ordinal sums of a copula* (Nelsen, 2006, p. 63) or *M-ordinal sum of the summands* (Klement *et al.*, 2017) within $\mathcal{I}^2$ into n partitions (possibly infinite) within $\mathcal{I}^2$. According to Nelsen, letting $\mathcal{J}$ denote a *partition* of $\mathcal{I}^2$ and $\mathcal{J}_i = [a_i, b_i]$ be the i th partition that does not overlap with others and letting also $\mathbf{C}_i$ be a copula for the i th partition, then the *ordinal sum* of these $\mathbf{C}_i$ with parameters Θ_i with respect to $\mathcal{J}_i$ is the copula $\mathbf{C}$ given by

$$\mathbf{C}(u, v; \mathcal{J}_i, \mathbf{C}_i, \Theta_i, i \in 1, 2, \dots, n) = a_i + (b_i - a_i) \mathbf{C}_i\left(\frac{u - a_i}{b_i - a_i}, \frac{v - a_i}{b_i - a_i}; \Theta_i\right) \text{ for } (u, v) \in \mathcal{J}_i,$$

for points within the partitions, and for points otherwise outside the partitions the coupla is given by

$$\mathbf{C}(u, v; \mathcal{J}_i, \mathbf{C}_i, i \in 1, 2, \dots, n) = \mathbf{M}(u, v) \text{ for } (u, v) \ni \mathcal{J}_i, \text{ and}$$

let $\mathbf{C}_{\mathcal{J}}(u, v)$ be a convenient abbreviation for the copula. Finally, Nelsen (2006, theorem 3.2.1) states that a copula is an ordinal sum if and only if for a t if $\mathbf{C}(t, t) = t$ for $t \in (0, 1)$. The *diagonal of a coupla* can be useful for quick assessment (see **Examples**) of this theorem. (See [ORDSUWcop](#), *W-ordinal sum of the summands.*)

Usage

```
ORDSUMcop(u,v, para=list(cop=W, para=NA, part=c(0,1)), ...)
```

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction;
para	A list of sublists for the coupla, parameters, and partitions (see Examples) and some attempt for intelligent in-fill of para is made within the sources (the default para is an example for which cop and para elements are converted to lists). The user is responsible that part element properly canvases by end-point alignment all of $\mathcal{I}^2$; and
...	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

Klement, E.P., Kolesárová, A., Mesiar, R., Saminger-Platz, S., 2017, Copula constructions using ultramodularity (chap. 9) in Copulas and dependence models with applications—Contributions in honor of Roger B. Nelsen, *eds.* Flores, U.M., Amo Artero, E., Durante, F., Sánchez, J.F.: Springer, Cham, Switzerland, ISBN 978–3–319–64220–9, doi:10.1007/9783319642215.

See Also

[copBasic-package, W_N5p12a, ORDSUMcop](#)

Examples

```
## Not run:
para <- list(cop=c(CLcop, M, PLcop, GHcop), para=list(4, NA, 0.1, c(3, 4)),
            part=list(c(0, 0.25), c(0.25, 0.35), c(0.35, 0.85), c(0.85, 1)))
UV <- simCOP(n=100, cop=ORDSUMcop, para=para, ploton=FALSE)
plot(c(0,1), c(0,1), xlab="U, nonexceedance probability", type="n", las=1,
     ylab="V, nonexceedance probability")
for(k in seq_len(length(para$part))) { # to draw the partitions
  a <- para$part[[k]][1]; b <- para$part[[k]][2]
  polygon(c(a, b, b, a, a), c(a, a, b, b, a), lty=2, lwd=0.8, col="lightgreen")
  text((a+b) / 2, (a+b) / 2, k, cex=3, col="blue") # numbered by partition
}
points(UV, pch=21, cex=0.8, col=grey(0.1), bg="white") #
## End(Not run)

## Not run:
para <- list(cop=c(GHcop), para=list(c(2, 3)), # internally replicated
            part=list(c(0, 0.2), c(0.2, 0.3), c(0.3, 0.5), c(0.5, 0.7), c(0.7, 1)))
UV <- simCOP(n=100, cop=ORDSUMcop, para=para, ploton=FALSE)
plot(c(0,1), c(0,1), xlab="U, nonexceedance probability", type="n", las=1,
     ylab="V, nonexceedance probability")
for(k in seq_len(length(para$part))) { # to draw the partitions
  a <- para$part[[k]][1]; b <- para$part[[k]][2]
  polygon(c(a, b, b, a, a), c(a, a, b, b, a), lty=2, lwd=0.8, col="lightgreen")
  text((a+b) / 2, (a+b) / 2, k, cex=3, col="blue") # numbered by partition
}
points(UV, pch=21, cex=0.8, col=grey(0.1), bg="white") #
## End(Not run)

## Not run:
# In this example, it is important that the delt is of the resolution
```

```

# matching the edges of the partitions.
para <- list(cop=P, para=list(NULL),
            part=list(c(0, 0.257), c(0.257, 0.358), c(0.358, 1)))
DI <- diagCOP(cop=ORDSUMcop, para=para, delt=0.001)
if(sum(DI$diagcop == DI$t) >= 1) {
  message("The ORDSUMcop() operation is an ordinal sum if there exists\n",
          "a t=(0,1) exists such that C(t,t)=t by Nelsen (2006, theorem 3.2.1).")
}
lines(par()$usr[1:2], par()$usr[3:4], col="red") #
## End(Not run)

## Not run:
# Expansive demonstration of high levels of "shape" of bivariate relations beyond
# the "simple" measures of association (Rho, Tau, Wolf, ...):
myCop <- ORDSUMcop # Note, integration problems more likely, if changed to
# ORDSUMcop at which point |Spearman Rhos| might integrate to beyond unity.
# Note, hardwired parameters in the demo-definitions of custom subcopulas to
# avoid confusion to study departure away from true M() and W() copulas, and to
# have these examples producing permutation asymmetry and L-comoments not exchangeable
# though only the with respect to UV [12] being plotted. We swap to M, W as shown
# in the example (three lines below) to make the default of example exchangeable.
myM <- function(u,v, para=NULL, ...) P(u, v, para=NULL, ...)
myW <- function(u,v, para=NULL, ...) u - PAcop(u, 1-v, para=7, ...) # rotate 90d
myM <- M; myW <- W # comment this line out for "custom subcopulas" after which
# the "myM" means first defined subcopula and "myW" means second defined subcopula.
# ----- Demonstrate with mid-partioning -----
mara <- list(cop=c(myM, myW), para=NULL, part=c(0, 0.5, 1))
UVm <- simCOP(1E3, cop=myCop, para=mara, seed=1, col="red")
wara <- list(cop=c(myW, myM), para=NULL, part=c(0, 0.5, 1))
UVw <- simCOP(1E3, cop=myCop, para=wara, seed=1, col="blue", ploton=FALSE)
cor(UVm[,1], UVm[,2], method="spearman"); cor(UVw[,1], UVw[,2], method="spearman")
# -----
opts <- par(las=1, lend=1, ljoin=1) # all lab horizontal, butted, mitered
del <- 0.005
nsim <- 300; ps <- c(0.001, seq(del, 1-del, by=del), 0.999)
# The ylim might need extension, it seems cases of L-comoments exceeding unity
# can be found depending on the settings of the myM and myW.
plot(c(0, 1), c(-1, 1), type="n", ylab="Spearman Rho, L-coskew, and L-cokurtosis",
     xlab="Ordinal sum partitioning parameter")
# -- Stuff below can be run again with changing the three logical
show_points <- FALSE # variables as user needs. We like TRUE
show_guidelines <- TRUE # FALSE, FALSE for a look by sampling
dolcomCOP <- TRUE # before FALSE, TRUE, TRUE for theory.
T2m12 <- T3m12 <- T4m12 <- T2w12 <- T3w12 <- T4w12 <- NULL # L-comoments
T2m21 <- T3m21 <- T4m21 <- T2w21 <- T3w21 <- T4w21 <- NULL # L-comoments
NuSk_m <- NuSt_m <- NuSk_w <- NuSt_w <- NULL # Joe "skews"
# lcomCOP() # theoretical L-comoments by integration
# lmomco:lmomoms2() # sample L-comoments estimation
# nuskewCOP() is a permutation asymmetry measure and nustarCOP() is something
# slightly more complex in what it measures, but it is important to note that
# each is in a linear combination to the dual integral of C(u,v), and therefore,
# we can expect them to be coupled to Spearman Rho itself in a rather simple way.
for(p in ps) { message(p, "-", appendLF=FALSE)

```

```

mara <- list(cop=c(myM, myW), para=NULL, part=c(0, p, 1))
wara <- list(cop=c(myW, myM), para=NULL, part=c(0, p, 1))
if(dolcomCOP) { # theoretical (numerical integration)
  lcm <- lcmCOP(cop=myCop, para=mara, stop.on.error=FALSE)
  lcw <- lcmCOP(cop=myCop, para=wara, stop.on.error=FALSE)
  T2m12 <- c(T2m12, lcm$lcomUV[2]); T2w12 <- c(T2w12, lcw$lcomUV[2])
  T2m21 <- c(T2m21, lcm$lcomVU[2]); T2w21 <- c(T2w21, lcw$lcomVU[2])
  T3m12 <- c(T3m12, lcm$lcomUV[3]); T4m12 <- c(T4m12, lcm$lcomUV[4])
  T3w12 <- c(T3w12, lcw$lcomUV[3]); T4w12 <- c(T4w12, lcw$lcomUV[4])
  T3m21 <- c(T3m21, lcm$lcomVU[3]); T4m21 <- c(T4m21, lcm$lcomVU[4])
  T3w21 <- c(T3w21, lcw$lcomVU[3]); T4w21 <- c(T4w21, lcw$lcomVU[4])
  NuSk_m <- c(NuSk_m, nuskeWCOP( cop=myCop, para=mara) )
  NuSt_m <- c(NuSt_m, nustarCOP( cop=myCop, para=mara) )
  NuSk_w <- c(NuSk_w, nuskeWCOP( cop=myCop, para=wara) )
  NuSt_w <- c(NuSt_w, nustarCOP( cop=myCop, para=wara) )
} else { # sample versions
  uvm <- rCOP(nsim, cop=myCop, para=mara, seed=1)
  uvw <- rCOP(nsim, cop=myCop, para=wara, seed=1)
  lcm <- lmomco::lcomoms2(uvm, nmom=4)
  lcm <- list(lcomUV=c(NA, lcm$T2[1,2], lcm$T3[1,2], lcm$T4[1,2]),
             lcomVU=c(NA, lcm$T2[2,1], lcm$T3[2,1], lcm$T4[2,1]))
  lcw <- lmomco::lcomoms2(uvw, nmom=4)
  lcw <- list(lcomUV=c(NA, lcw$T2[1,2], lcw$T3[1,2], lcw$T4[1,2]),
             lcomVU=c(NA, lcw$T2[2,1], lcw$T3[2,1], lcw$T4[2,1]))
  T2m12 <- c(T2m12, lcm$lcomUV[2]); T2w12 <- c(T2w12, lcw$lcomUV[2])
  T2m21 <- c(T2m21, lcm$lcomVU[2]); T2w21 <- c(T2w21, lcw$lcomVU[2])
  T3m12 <- c(T3m12, lcm$lcomUV[3]); T4m12 <- c(T4m12, lcm$lcomUV[4])
  T3w12 <- c(T3w12, lcw$lcomUV[3]); T4w12 <- c(T4w12, lcw$lcomUV[4])
  T3m21 <- c(T3m21, lcm$lcomVU[3]); T4m21 <- c(T4m21, lcm$lcomVU[4])
  T3w21 <- c(T3w21, lcw$lcomVU[3]); T4w21 <- c(T4w21, lcw$lcomVU[4])
  NuSk_m <- c(NuSk_m, nuskeWCOP( para=uvm, as.sample=TRUE) )
  NuSt_m <- c(NuSt_m, nustarCOP( para=uvm, as.sample=TRUE) )
  NuSk_w <- c(NuSk_w, nuskeWCOP( para=uvw, as.sample=TRUE) )
  NuSt_w <- c(NuSt_w, nustarCOP( para=uvw, as.sample=TRUE) )
}
# -----
if(show_points) {
  n <- length(T3m12)
  points(p, T2m12[n], pch="R", col="seagreen4" ) # Spearman Rho
  points(p, T2w12[n], pch="R", col="darkorchid4" ) # Spearman Rho
  points(p, T3m12[n], pch=24, col="seagreen4", bg="seagreen4" )
  points(p, T4m12[n], pch=25, col="seagreen4", bg="seagreen4" )
  points(p, T3w12[n], pch=24, col="darkorchid4", bg="darkorchid4")
  points(p, T4w12[n], pch=25, col="darkorchid4", bg="darkorchid4")
  points(p, NuSk_m[n], pch=2, col="deepskyblue2") # open upright tri
  points(p, NuSt_m[n], pch=8, col="seagreen4" ) # pch == star
  points(p, NuSk_w[n], pch=6, col="deepskyblue2") # open flipped tri
  points(p, NuSt_w[n], pch=8, col="darkorchid4" ) # pch == star
}
}
message("done")
if(show_guidelines) {
  psna <- c(ps, NA, ps) # insert NA to "line break"
}

```

```

lines(psna, c( T2m12, NA, T2w12), lwd=3, col="grey80")
lines(psna, c( T3m12, NA, T3w12), lty=1)
lines(psna, c( T4m12, NA, T4w12), lty=2)
lines(psna, c(NuSk_m, NA, NuSk_w), lty=1)
lines(psna, c(NuSt_m, NA, NuSt_w), lty=2)
}
par(opts)
# Note, the nuskews will plot along origin line if the "image" of the two
# subcouplas are permutation symmetric and result in a hexagram (six-
# sided star) being "plotted." If subcopulas themselves are not permutation
# symmetric, then triangles for nuskew will diverge from each other.
## End(Not run)

## Not run:
# Now, look at the coupling between the Spearman Rho and the L-comoments through
# formation of L-comoment ratio diagrams of the ordinal sums.
# Here, we just draw lines from whatever the vectors stemming from the previous
# example produced by dolcomCOP being set TRUE (theoretical) or FALSE (sample)
# L-comoments. For the simple, ordinal sum of M,W or W,M, then L-cokurtosis
# will plot on top of each other; therefore, we use the thick line for T4m12.
plot(c(-1, 1), c(-1, 1), type="n", las=1,
      xlab="Spearman Rho (driven by ordinal sum parition)"
      ylab="L-coskew (cT3) or L-cokurtosis (cT4)")
lines(par()$usr[1:2], rep(0, 2))
lines(T2m12, T3m12, lty=1, col="seagreen4", lwd=3)
lines(T2m12, T4m12, lty=2, col="seagreen4", lwd=3)
lines(T2w12, T3w12, lty=1, col="darkorchid4")
lines(T2w12, T4w12, lty=2, col="darkorchid4") # (myM,myW);(myW,myM) plot on T4m21?
txt <- c("L-coskew cT3[12], cop(myM, myW)", "L-cokurtosis cT4[12], cop(myM, myW)",
        "L-coskew cT3[12], cop(myW, myM)", "L-cokurtosis cT4[12], cop(myW, myM)")
legend("topleft", txt, bty="n", seg.len=3, lty=c(1, 2, 1, 2), lwd=c(3, 3, 1, 1),
      col=c("seagreen4", "seagreen4", "darkorchid4", "darkorchid4"))
# -----
# If the user is working with sufficiently complex copulas, then differing
# trajectories through the domain occur.
plot(c(-1, 1), c(-1, 1), type="n", las=1, ylab="L-coskew or L-cokurtosis",
      xlab="Spearman Rho (driven by ordinal sum partition)")
lines(par()$usr[1:2], rep(0, 2))
lines(T2m21, T3m21, lty=1, col="seagreen4", lwd=3)
lines(T2m21, T4m21, lty=2, col="seagreen4", lwd=3)
lines(T2w21, T3w21, lty=1, col="darkorchid4")
lines(T2w21, T4w21, lty=2, col="darkorchid4") # (myM,myW);(myW,myM) plot on T4m21?
txt <- c("L-coskew cT3[21], cop(myM, myW)", "L-cokurtosis cT4[21], cop(myM, myW)",
        "L-coskew cT3[21], cop(myW, myM)", "L-cokurtosis cT4[21], cop(myW, myM)")
legend("topleft", txt, bty="n", seg.len=3, lty=c(1, 2, 1, 2), lwd=c(3, 3, 1, 1),
      col=c("seagreen4", "seagreen4", "darkorchid4", "darkorchid4")) #
## End(Not run)

## Not run:
# -----
# Next, let us build polynomial approximations for the boundaries and the polynomial
# degree has been set pretty large in order that the user would have built-in
# flexibility to construction of these approximations for more complex copula than

```

```

# the stacking of M,W and W,M.
X   <- c(T2m12, T2m21); Y3mw <- c(T3m12, T3m21); xs <- seq(-1,1, by=0.001)
T3fT2mw <- lm(Y3mw~I(X^ 1)+I(X^ 2)+I(X^ 3)+I(X^ 4)+I(X^ 5)+I(X^ 6)+
               I(X^ 7)+I(X^ 8)+I(X^ 9)+I(X^10)+I(X^11)+I(X^12))
plot(X, Y3mw, xlab="Spearman Rho", ylab="L-coskew (cT3)")
lines(xs, predict(T3fT2mw, newdata=data.frame(X=xs)), col="red")
mtext("2-partition ORDSUMcop : M, W")

X   <- c(T2w12, T2w21); Y3wm <- c(T3w12, T3w21)
T3fT2wm <- lm(Y3wm~I(X^ 1)+I(X^ 2)+I(X^ 3)+I(X^ 4)+I(X^ 5)+I(X^ 6)+
               I(X^ 7)+I(X^ 8)+I(X^ 9)+I(X^10)+I(X^11)+I(X^12))
plot(X, Y3wm, xlab="Spearman Rho", ylab="L-coskew (cT3)")
lines(xs, predict(T3fT2wm, newdata=data.frame(X=xs)), col="red")
mtext("2-partition ORDSUMcop : W, M")

X   <- c(T2m12, T2m21, T2w12, T2w21); Y4 <- c(T4m12, T4m21, T4w12, T4w21)
T4fT2mwwm <- lm(Y4~I(X^ 1)+I(X^ 2)+I(X^ 3)+I(X^ 4)+I(X^ 5)+I(X^ 6)+
                 I(X^ 7)+I(X^ 8)+I(X^ 9)+I(X^10)+I(X^11)+I(X^12))

plot(X, Y4, xlab="Spearman Rho", ylab="L-cokurtosis (cT4)")
lines(xs, predict(T4fT2mwwm, newdata=data.frame(X=xs)), col="red")

X   <- c(T3m12, T3m21, T3w12, T3w21)
Y4  <- c(T4m12, T4m21, T4w12, T4w21)
plot(X, Y4, type="l", xlab="L-coskew", ylab="L-cokurtosis")
points(T3m12, T4m12, col="seagreen4")
points(T3w12, T4w12, col="orchid3" ) #
## End(Not run)

## Not run:
# -----
# Next, let us extract the coefficients and build-up stand alone functions that
# could be used to draw the L-comoments of the ordinal sums and potentially
# overlaid on L-comoment ratio diagrams such as seen built under the Examples in the
# LCOMDIA*.Rd documentation files (for instance LCOMDIA_ManyCops.Rd) of the package.
coesA <- T3fT2mw$coefficients; names(coesA) <- NULL
coesB <- T3fT2wm$coefficients; names(coesB) <- NULL
coesC <- T4fT2mwwm$coefficients; names(coesC) <- NULL
print(coesA, 12); print(coesB, 12); print(coesC, 12)
coesT3 <- sapply(seq_len(length(coesA)), function(i) {
  mean(c(abs(coesA[i]), abs(coesB[i]))) })
print(coesT3, 12)
"T34fT2" <- function(rho, type=c("T3mw", "T3wm", "T4mwwm")) {
  type <- match.arg(type)
  cs3 <- c(0.618673707079,
           0.175958850204, 0.512576942435, 0.145431255262, 0.265479078036,
           0.163312255777, 1.016327127995, 0.537153927577, 2.225271502140,
           0.636080402864, 2.191529289514, 0.292183399458, 0.823239656100)
  if(type == "T3mw") {
    cs <- c(-1,-1, 1, 1, 1,-1,-1, 1, 1,-1,-1, 1, 1)*cs3
  } else if(type == "T3wm") {
    cs <- c( 1, 1,-1,-1,-1, 1, 1,-1,-1, 1, 1,-1,-1)*cs3
  } else if(type == "T4mwwm") {

```

```

cs <- c( 0.236681925424,
        -0.429191591357, -0.573421581827, 0.352083105242, 1.042508365628,
        -0.710792468014, -4.321529301154, 2.229039543342, 9.419379550877,
        -2.652104491697, -9.275892279321, 1.208446517629, 3.472344853036)
snmw <- rep(1, length(cs))
} else {
  stop("should not be here in logic")
}
z <- sapply(rho, function(r) sum(sapply(seq_len(length(cs)),
                                       function(p) cs[p] * r^(p-1))))

return(z)
}

# Finally, plot again the L-comoment ratio diagram of Spearman Rho and the
# two higher L-comoments and a L-coskew and L-cokurtosis diagram could also be
# readily built from a vector of Spearman Rhos.
ylim <- c(-0.7, 0.7)
plot( xs, T34fT2(xs, "T3mw"), type="n", ylim=ylim,
      xlab="Spearman Rho", ylab="L-coskew or L-cokurtosis approximations")
lines(par()$usr[1:2], rep(0, 2))
lines(xs, T34fT2(xs, "T3mw"), col="brown3")
lines(xs, T34fT2(xs, "T3wm"), col="blue3" )
lines(xs, T34fT2(xs, "T4mwwm"), col="orange")
# -----
ylim <- c(-0.7, 0.7)
plot(c(T34fT2(xs, "T3mw" ), T34fT2(xs, "T3wm" )),
     c(T34fT2(xs, "T4mwwm"), T34fT2(xs, "T4mwwm")), type="n", ylim=ylim,
     xlab="L-coskew approximations", ylab="L-cokurtosis approximations")
lines(par()$usr[1:2], rep(0, 2)); lines(rep(0, 2), par()$usr[3:4])
lines(T34fT2(xs, "T3mw"), T34fT2(xs, "T4mwwm"), col="seagreen4")
lines(T34fT2(xs, "T3wm"), T34fT2(xs, "T4mwwm"), col="orchid3" )
text(-0.35, 0.07, "Ordinal sum of\nM and W", font=2, col="seagreen4")
text(+0.35, 0.07, "Ordinal sum of\nW and M", font=2, col="orchid3" ) #
## End(Not run)

```

Description

Compute *W-ordinal sum of the summands* (Klement *et al.*, 2017) within $\mathcal{I}^2$ into n partitions (possibly infinite) within $\mathcal{I}^2$. Letting $\mathcal{J}$ denote a *partition* of $\mathcal{I}^2$ and $\mathcal{J}_i = [a_i, b_i]$ be the i th partition that does not overlap with others and letting also C_i be a copula for the i th partition, then the *ordinal sum* of these C_i with parameters Θ_i with respect to $\mathcal{J}_i$ is the copula C given by

$$C(u, v; \mathcal{J}_i, C_i, \Theta_i, i \in 1, 2, \dots, n) = a_i + (b_i - a_i) C_i \left(\frac{u - a_i}{b_i - a_i}, \frac{v - 1 + b_i}{b_i - a_i}; \Theta_i \right) \text{ for } (u, v) \in \mathcal{J}_i,$$

for points within the partitions, and for points otherwise outside the partitions the coupla is given by

$$\mathbf{C}(u, v; \mathcal{J}_i, \mathbf{C}_i, i \in 1, 2, \dots, n) = \mathbf{W}(u, v) \text{ for } (u, v) \ni \mathcal{J}^2, \text{ and}$$

let $\mathbf{C}_{\mathcal{J}}(u, v)$ be a convenient abbreviation for the copula. (See [ORDSUMcop](#), *M-ordinal sum of the summands*.)

Usage

```
ORDSUWcop(u,v, para=list(cop=M, para=NA, part=c(0,1)), ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A list of sublists for the coupla, parameters, and partitions (see Examples) and some attempt for intelligent in-fill of <code>para</code> is made within the sources (the default <code>para</code> is an example for which <code>cop</code> and <code>para</code> elements are converted to lists). The user is responsible that <code>part</code> element properly canvases by end-point alignment all of $\mathcal{I}^2$; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Klement, E.P., Kolesárová, A., Mesiar, R., Saminger-Platz, S., 2017, Copula constructions using ultramodularity (chap. 9) in *Copulas and dependence models with applications—Contributions in honor of Roger B. Nelsen*, eds. Flores, U.M., Amo Artero, E., Durante, F., Sánchez, J.F.: Springer, Cham, Switzerland, ISBN 978–3–319–64220–9, [doi:10.1007/9783319642215](https://doi.org/10.1007/9783319642215).

See Also

[copBasic-package](#), [W_N5p12a](#), [ORDSUMcop](#)

Examples

```
para <- list(cop=c(CLcop, GHcop), para=list(5, 2), part=c(0,0.25,1)) # break points
UV <- simCOP(n=100, cop=ORDSUMcop, seed=1, para=para, ploton=TRUE, pch=16)
UV <- simCOP(n=100, cop=ORDSUWcop, seed=1, para=para, ploton=FALSE)

## Not run:
para <- list(cop=c(CLcop, M, PLcop, GHcop), para=list(4, NA, 0.1, c(3, 4)),
             part=list(c(0, 0.25), c(0.25, 0.35), c(0.35, 0.85), c(0.85, 1)))
UV <- simCOP(n=100, cop=ORDSUWcop, para=para, ploton=FALSE)
```

```

plot(c(0,1), c(0,1), xlab="U, NONEXCEEDANCE PROBABILITY", type="n", las=1,
     ylab="V, NONEXCEEDANCE PROBABILITY")
for(k in seq_len(length(para$part))) {      # to draw the partitions
  a <- para$part[[k]][1]; b <- para$part[[k]][2]
  polygon(c(a, b, b, a, a), c(1-a,1-a,1-b,1-b,1-a), lty=2, lwd=0.8, col="lightgreen")
  text( (a+b) / 2, (1-a+1-b) / 2, k, cex=3, col="blue") # numbered by partition
}
points(UV, pch=21, cex=0.8, col=grey(0.1), bg="white") #
## End(Not run)

## Not run:
para = list(cop=c(GHcop), para=list(c(2, 3)), # internally replicated
            part=list(c(0, 0.2), c(0.2, 0.3), c(0.3, 0.5), c(0.5, 0.7), c(0.7, 1)))
UV <- simCOP(n=100, cop=ORDSUWcop, para=para, ploton=FALSE)
plot(c(0,1), c(0,1), xlab="U, NONEXCEEDANCE PROBABILITY", type="n", las=1,
     ylab="V, NONEXCEEDANCE PROBABILITY")
for(k in seq_len(length(para$part))) {      # to draw the partitions
  a <- para$part[[k]][1]; b <- para$part[[k]][2]
  polygon(c(a, b, b, a, a), c(1-a,1-a,1-b,1-b,1-a), lty=2, lwd=0.8, col="lightgreen")
  text( (a+b) / 2, (1-a+1-b) / 2, k, cex=3, col="blue") # numbered by partition
}
points(UV, pch=21, cex=0.8, col=grey(0.1), bg="white") #
## End(Not run)

```

P

The Product (Independence) Copula

Description

Compute the *product copula* (Nelsen, 2006, p. 12), which is defined as

$$\Pi(u, v) = uv.$$

This is the copula of statistical independence between U and V and is sometimes referred to as the *independence copula*. The two extreme antithesis copulas are the *Fréchet–Hoeffding upper-bound* ([M](#)) and *Fréchet–Hoeffding lower-bound* ([W](#)) copulas.

Usage

```
P(u, v, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[M](#), [W](#), [rhoCOP](#), [lcomCOP](#), [PLcop](#)

Examples

```
P(c(0.4, 0, 1), c(0, 0.6, 1))

## Not run:
# PERFECT INDEPENDENCE
n <- 100000 # large sample size, L-comoments are zero
lcomCOP(cop=P) # zeros (numerical integration)
UV <- simCOP(n=n, cop=P, graphics=FALSE)
lmomco::lcomoms2(UV, nmom=4)
# The following are Taus_r^{12} and Taus_r^{21}
# L-corr:      0.00265 and 0.00264 ---> ZERO
# L-coskew:    -0.00121 and 0.00359 ---> ZERO
# L-cokurtosis: 0.00123 and 0.00262 ---> ZERO

# CONTRAST TO MODEST POSITIVE CORRELATION
rho <- 0.6; # Spearman Rho
theta <- PLACKETTpar(rho=rho) # Theta = 5.115658
lcomCOP(cop=PLcop, para=theta) # Rho = 0.6 and rest zeros
UV <- simCOP(n=n, cop=PLcop, para=theta, graphics=FALSE)
lmomco::lcomoms2(UV, nmom=4)
# The following are Taus_r^{12} and Taus_r^{21}
# L-corr      0.60303 and 0.60303 ---> Spearman Rho
# L-coskews    4.5339e-05 and -0.00124011 ---> ZERO
# L-cokurtosis 0.00088549 and 0.00129812 ---> ZERO
# Pearson R == Spearman Rho for UV L-comoments
## End(Not run)
```

PARETOcop

The Pareto Copula

Description

The *Pareto copula* (Nelsen, 2006, pp. 33) is

$$C_{\Theta}(u, v) = \mathbf{PA}(u, v) = [(1 - u)^{-\Theta} + (1 - v)^{-\Theta}]^{-1/\Theta},$$

where $\Theta \in [0, \infty)$. As $\Theta \rightarrow 0^+$, the copula limits to the Π copula ([P](#)) and the $\mathbf{M}$ copula ([M](#)). The parameterization here has association increasing with increasing Θ , which differs from Nelsen

(2006), and also the Pareto copula is formed with right-tail increasing reflection of the Nelsen (2006) presentation because it is anticipated that **copBasic** users are more likely to have right-tail dependency situations (say large maxima [right tail] coupling in earth-system data but not small maxima [left tail] coupling).

Usage

```
PARETOcop(u, v, para=NULL, ...)
PAcop(u, v, para=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Note

The Pareto copula is used in a demonstration of *Kendall Function L-moment ratio diagram* construction (see [kfuncCOPlmoms](#)).

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[M](#), [P](#)

Examples

```
## Not run:
z <- seq(0.01, 0.99, by=0.01) # Both copulas have Kendall Tau = 1/3
plot( z, kfuncCOP(z, cop=PAcop, para=1), lwd=2, col="black",
      xlab="z <= Z", ylab="F_K(z)", type="l")
lines(z, kfuncCOP(z, cop=GHCop, para=1.5), lwd=2, col="red") # red line
# All extreme value copulas have the same Kendall Function [F_K(z)], the
# Gumbel-Hougaard is such a copula and the F_K(z) for the Pareto does not
# plot on top and thus is not an extreme value but shares a "closeness."
## End(Not run)
```

Description

The *Plackett copula* (Nelsen, 2006, pp. 89–92) is

$$\mathbf{C}_\Theta(u, v) = \mathbf{PL}(u, v) = \frac{[1 + (\Theta - 1)(u + v)] - \sqrt{[1 + (\Theta - 1)(u + v)]^2 - 4uv\Theta(\Theta - 1)}}{2(\Theta - 1)}.$$

The Plackett copula ($\mathbf{PL}(u, v)$) is *comprehensive* because as $\Theta \rightarrow 0$ the copula becomes $\mathbf{W}(u, v)$ (see [W](#), *countermonotonicity*), as $\Theta \rightarrow \infty$ the copula becomes $\mathbf{M}(u, v)$ (see [M](#), *comonotonicity*) and for $\Theta = 1$ the copula is $\mathbf{\Pi}(u, v)$ (see [P](#), *independence*).

Nelsen (2006, p. 90) shows that

$$\Theta = \frac{H(x, y)[1 - F(x) - G(y) + H(x, y)]}{[F(x) - H(x, y)][G(y) - H(x, y)]},$$

where $F(x)$ and $G(y)$ are cumulative distribution function for random variables X and Y , respectively, and $H(x, y)$ is the joint distribution function. Only Plackett copulas have a constant Θ for any pair $\{x, y\}$. Hence, Plackett copulas are also known as *constant global cross ratio* or *contingency-type* distributions. The copula therefore is intimately tied to *contingency tables* and in particular the bivariate Plackett defined herein is tied to a 2×2 contingency table. Consider the 2×2 contingency table shown at the end of this section, then Θ is defined as

$$\Theta = \frac{a/c}{b/d} = \frac{\frac{a}{a+c}/\frac{c}{a+c}}{\frac{b}{b+d}/\frac{d}{b+d}} \text{ and } \Theta = \frac{a/b}{c/d} = \frac{\frac{a}{a+b}/\frac{b}{a+b}}{\frac{c}{c+d}/\frac{d}{c+d}},$$

where it is obvious that $\Theta = ad/bc$ and a, b, c , and d can be replaced by proportions for a sample of size n by $a/n, b/n, c/n$, and d/n , respectively. Finally, this copula has been widely used in modeling and as an alternative to bivariate distributions and has respective *lower-* and *upper-tail dependency parameters* of $\lambda^L = 0$ and $\lambda^U = 0$ ([taildepCOP](#)).

--	Low	High	Sums
Low	a	b	$a + b$
High	c	d	$c + d$
Sums	$a + c$	$b + d$	--

Usage

```
PLACKETTcop(u, v, para=NULL, ...)
PLcop(u, v, para=NULL, ...)
```

Arguments

u Nonexceedance probability u in the X direction;
v Nonexceedance probability v in the Y direction;
para A vector (single element) of parameters—the Θ parameter of the copula; and
... Additional arguments to pass.

Value

Value(s) for the copula are returned.

Note

The Plackett copula was the first (2008) copula implemented in **copBasic** as part of initial development of the code base for instructional purposes. Thus, this particular copula has a separate parameter estimation function in [PLACKETTpar](#) as a historical vestige of a class project.

Author(s)

W.H. Asquith

References

- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
 Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[PLACKETTpar](#), [PLpar](#), [PLACKETTsim](#), [W](#), [M](#), [densityCOP](#)

Examples

```
PLACKETTcop(0.4, 0.6, para=1)
P(0.4, 0.6) # independence copula, same two values because Theta == 1
PLcop(0.4, 0.6, para=10.25) # joint probability through positive association

## Not run:
# Joe (2014, p. 164) shows the closed form copula density of the Plackett.
"dPLACKETTcop" <- function(u,v,para) {
  eta <- para - 1; A <- para*(1 + eta*(u+v-2*u*v))
  B <- ((1 + eta*(u+v))^2 - 4*para*eta*u*v)^(3/2); return(A/B)
}
u <- 0.08; v <- 0.67 # Two probabilities to make numerical evaluations.
del <- 0.0001 # a 'small' differential value of probability
u1 <- u; u2 <- u+del; v1 <- v; v2 <- v+del
# Density following (Nelsen, 2006, p. 10)
dCrect <- (PLcop(u2, v2, para=10.25) - PLcop(u2, v1, para=10.25) -
  PLcop(u1, v2, para=10.25) + PLcop(u1, v1, para=10.25)) / del^2
dCanal <- dPLACKETTcop(u, v, para=10.25)
dCfunc <- densityCOP(u, v, para=10.25, cop=PLcop, deluv = del)
R <- round(c(dCrect, dCanal, dCfunc), digits=6)
message("Density: ", R[1], "(manual)", ", R[2], "(analytical)", ", R[3], "(function)");
# Density: 0.255377(manual), 0.255373(analytical), 0.255377(function)

# Comparison of partial derivatives
dUr <- (PLcop(u2, v2, para=10.25) - PLcop(u1, v2, para=10.25)) / del
dVr <- (PLcop(u2, v2, para=10.25) - PLcop(u2, v1, para=10.25)) / del
dU <- derCOP( u, v, cop=PLcop, para=10.25)
dV <- derCOP2(u, v, cop=PLcop, para=10.25)
```

```
R <- round(c(dU, dV, dUr, dVr), digits=6)
message("Partial derivatives dU=", R[1], " and dUr=", R[3], "\n",
"      "      "dV=", R[2], " and dVr=", R[4]) #
## End(Not run)
```

PLACKETTpar

Estimate the Parameter of the Plackett Copula

Description

The parameter Θ of the *Plackett copula* (Nelsen, 2006, pp. 89–92) ([PLACKETTcop](#) or [PLcop](#)) is related to the *Spearman Rho* ($\rho_S \neq 1$, see [rhoCOP](#))

$$\rho_S(\Theta) = \frac{\Theta + 1}{\Theta - 1} - \frac{2\Theta \log(\Theta)}{(\Theta - 1)^2}.$$

Alternatively, the parameter can be estimated using a *median-split estimator*, which is best shown as an algorithm. First, compute the two medians:

```
medx <- median(x); medy <- median(y)
```

Second and third, compute the number of occurrences where both values are less than their medians and express that as a probability:

```
k <- length(x[x < medx & y < medy]); m <- k / length(x)
```

Finally, the median-split estimator of Θ is computed by

$$\Theta = \frac{4m^2}{(1 - 2m)^2}.$$

Nelsen (2006, p. 92) and Salvadori *et al.* (2007, p. 247) provide further details. The input values x and y are *not used* for the median-split estimator if *Spearman Rho* (see [rhoCOP](#)) is provided by `rho`.

Usage

```
PLACKETTpar(x, y, rho=NULL, byrho=FALSE, cor=NULL, ...)
PLpar(x, y, rho=NULL, byrho=FALSE, cor=NULL, ...)
```

Arguments

<code>x</code>	Vector of values for random variable X ;
<code>y</code>	Vector of values for random variable Y ;
<code>rho</code>	Spearman Rho and <code>byrho</code> is set to TRUE automatically;
<code>byrho</code>	Should Spearman Rho be used instead of the median-split estimator;
<code>cor</code>	A copBasic syntax for “the correlation coefficient” suitable for the copula—a synonym for <code>rho</code> ; and
<code>...</code>	Additional arguments to pass.

Value

A value for the Plackett copula Θ is returned.

Note

Evidently there “does not appear to be a closed form for $\tau(\Theta)$ ” (Fredricks and Nelsen, 2007, p. 2147), but given $\rho(\Theta)$, the equivalent $\tau(\Theta)$ can be computed by either the [tauCOP](#) function or by approximation. One of the Examples sweeps through $\rho \mapsto [0, 1; \Delta\rho=\delta]$, fits the Plackett $\theta(\rho)$, and then solves for Kendall Tau $\tau(\theta)$ using [tauCOP](#). A polynomial is then fit between τ and ρ to provide rapid conversion between $|\rho|$ and τ , where the residual standard error is 0.0005705, adjusted R-squared is ≈ 1 , the maximum residual is $\epsilon < 0.006$. Because of symmetry, it is only necessary to fit positive association and reflect the result by the sign of ρ . This polynomial is from the Examples is

```
rho <- 0.920698
"getPLACKETTtau" <- function(rho) {
  taupoly <- 0.6229945*abs(rho) + 1.1621854*abs(rho)^2 -
    10.7424188*abs(rho)^3 + 48.9687845*abs(rho)^4 -
    119.0640264*abs(rho)^5 + 160.0438496*abs(rho)^6 -
    111.8403591*abs(rho)^7 + 31.8054602*abs(rho)^8
  return(sign(rho)*taupoly)
}
getPLACKETTtau(rho) # 0.7777726
```

The following code might be useful in some applications for the inversion of the polynomial for the ρ as a function of τ :

```
"fun" <- function(rho, tau=NULL) {tp <- getPLACKETTtau(rho); return(tau-tp)}
tau <- 0.78
rho <- uniroot(fun, interval=c(0, 1), tau=tau)$root # 0.9220636
```

Author(s)

W.H. Asquith

References

- Fredricks, G.A. and Nelsen, R.B., 2007, On the relationship between Spearman's rho and Kendall's tau for pairs of continuous random variables: *Journal of Statistical Planning and Inference*, v. 137, pp. 2143–2150.
- Nelsen, R.B., 2006, *An introduction to copulas*: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, *Extremes in Nature—An approach using copulas*: Springer, 289 p.

See Also

[PLACKETTcop](#), [PLcop](#), [PLACKETTsim](#), [rhoCOP](#)

Examples

```
## Not run:
Q1 <- rnorm(1000); Q2 <- Q1 + rnorm(1000)
PLpar(Q1, Q2); PLpar(Q1, Q2, byrho=TRUE) # two estimates for same data
PLpar(rho= 0.76) # positive association
PLpar(rho=-0.76) # negative association
tauCOP(cop=PLcop, para=PLpar(rho=-0.15, by.rho=TRUE)) #
## End(Not run)

## Not run:
RHOS <- seq(0, 0.990, by=0.002); TAUS <- rep(NA, length(RHOS))
for(i in 1:length(RHOS)) {
  #message("Spearman Rho: ", RHOS[i])
  theta <- PLACKETTpar(rho=RHOS[i], by.rho=TRUE); tau <- NA
  try(tau <- tauCOP(cop=PLACKETTcop, para=theta), silent=TRUE)
  TAUS[i] <- ifelse(is.null(tau), NA, tau)
}
LM <- lm(TAUS~ RHOS + I(RHOS^2) + I(RHOS^3) + I(RHOS^4) +
          I(RHOS^5) + I(RHOS^6) + I(RHOS^7) + I(RHOS^8) - 1)
plot(RHOS,TAUS, type="l", xlab="abs(Spearman Rho)", ylab="abs(Kendall Tau)")
lines(RHOS,fitted.values(LM), col=3)#
## End(Not run)
```

 PLACKETTsim

Direct Simulation of a Plackett Copula

Description

Plackett copula simulation (Nelsen, 2006, pp. 89–92) ([PLACKETTcop](#)) is made by this function using analytical formula (Durante, 2007, p. 247; see source code). The PLACKETTsim function exists for comparison against the numerical derivative (*conditional distribution method*) methods ([simCOP](#), [simCOPmicro](#)) otherwise used in **copBasic**.

Usage

```
PLACKETTsim(n, para=NULL, ...)
```

Arguments

n	Sample size;
para	The Θ parameter of the Plackett copula; and
...	Additional arguments to pass.

Value

An R data.frame of the values U and V for the nonexceedance probabilities is returned.

Author(s)

W.H. Asquith

References

Durante, F., 2007, Families of copulas, Appendix C, *in* Salvadori, G., De Michele, C., Kotegoda, N.T., and Rosso, R., 2007, Extremes in Nature—An approach using copulas: Springer, 289 p.

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also[PLACKETTcop](#), [PLACKETTpar](#)**Examples**

```
PLACKETTsim(10, para= 1 ) # simulate P (independence) copula through a Plackett
PLACKETTsim(10, para=20.3) # simulate strong positive Plackett
```

prod2COP

*The Product of Two Copulas***Description**

Perform copula multiplication (so-called “*-product” or *Markov Product*) (Darsow and others, 1992) is a continuous analog of matrix multiplication and yields another copula:

$$(\mathbf{C}_1 * \mathbf{C}_2)(u, v) = \mathbf{C}_3(u, v) = \int_{\mathcal{I}} \frac{\delta \mathbf{C}_1(u, t)}{\delta v} \frac{\delta \mathbf{C}_2(t, v)}{\delta u} dt,$$

for copulas $\mathbf{C}_1(u, v)$ and $\mathbf{C}_2(u, v)$ are copulas whose *-product yields copula $\mathbf{C}_3(u, v)$ in terms of partial derivatives ([derCOP](#) and [derCOP2](#)) of the other two. Nelsen (2006, p. 245) lists several identities of the *-product involving the product ($\mathbf{\Pi}$; $\mathbf{P}$), lower bound ($\mathbf{W}$; $\mathbf{w}$), and upper bound ($\mathbf{M}$; $\mathbf{m}$) copulas:

$$\mathbf{\Pi} * \mathbf{C} = \mathbf{C} * \mathbf{\Pi} = \mathbf{\Pi},$$

$$\mathbf{M} * \mathbf{C} = \mathbf{C} * \mathbf{M} = \mathbf{M},$$

$$(\mathbf{W} * \mathbf{C})(u, v) = v - \mathbf{C}(1 - u, v) \text{ and } (\mathbf{C} * \mathbf{W})(u, v) = u - \mathbf{C}(u, 1 - v), \text{ and}$$

$$\mathbf{W} * \mathbf{W} = \mathbf{M} \text{ and } \mathbf{W} * \mathbf{C} * \mathbf{W} = \hat{\mathbf{C}},$$

where $\hat{\mathbf{C}}$ is the *survival copula* ([surCOP](#)). The *-product is associative:

$$\mathbf{A} * (\mathbf{B} * \mathbf{C}) = (\mathbf{A} * \mathbf{B}) * \mathbf{C},$$

but *-product is not commutative (order independent). Nelsen (2006, p. 245) reports that “if we view * as a binary operation on the set of copulas, then $\mathbf{\Pi}$ is the null element, and $\mathbf{M}$ is the identity.” Copula multiplication is closely linked to *Markov Processes* (Nelsen, 2006, pp. 244–248).

For other descriptions and computations of copula combination are possible using the **copBasic** package, see [convexCOP](#), [convex2COP](#), [composite1COP](#), [composite2COP](#), [composite3COP](#), [glueCOP](#), and [convexCOP](#).

Usage

```
prod2COP(u,v, cop1=NULL, para1=NULL, cop2=NULL, para2=NULL, para=NULL,
          pinterval=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>cop1</code>	The $C_1(u, v; \Theta_1)$ copula function with vectorization as in <code>asCOP</code> ;
<code>para1</code>	Vector of parameters or other data structures for Θ_1 , if needed, to pass to copula $C_1(u, v; \Theta_1)$;
<code>cop2</code>	The $C_2(u, v; \Theta_2)$ copula function with vectorization as in <code>asCOP</code> ;
<code>para2</code>	Vector of parameters or other data structures for Θ_2 , if needed, to pass to copula $C_2(u, v; \Theta_2)$;
<code>para</code>	An R list that can take the place of the <code>cop1</code> , <code>para1</code> , <code>cop2</code> , and <code>para2</code> arguments. These four will be populated from same named elements of the list, and if the other four arguments were specified through the function interface, these are silently ignored;
<code>pinterval</code>	An optional interval for the above integral. The default is $\mathcal{I} = [0, 1]$ but the option of the user to replace exact end points with “small” numbers is possible (e.g. <code>interval=c(lo, 1-lo)</code> for say <code>lo=.Machine\$double.eps</code>). This interval is uniquely picked up for the interval in the above definition of <code>prod2COP</code> . The <code>pinterval</code> can also be set within the <code>para</code> and the function will pick it up from there; and
<code>...</code>	Additional arguments to pass to the copulas.

Value

Value(s) for the copula are returned.

Note

The *Farlie–Gumbel–Morgenstern copula* ($\mathbf{FGM}(u, v; \Theta)$; `FGMcop`) is

$$\mathbf{FGM}(u, v; \Theta) = uv[1 + \Theta(1 - u)(1 - v)],$$

where $-1 \leq \Theta \leq 1$. Nelsen (2006, exer. 6.12, p. 249) asserts that for $\mathbf{FGM}_{(\Theta=\alpha)}$ and $\mathbf{FGM}_{(\Theta=\beta)}$ with $*$ -product as $\mathbf{FGM}_\alpha * \mathbf{FGM}_\beta$ that a closed-form solution exists and is

$$\mathbf{FGM}_\alpha * \mathbf{FGM}_\beta = \mathbf{FGM}_{(\alpha\beta)/3}.$$

This assertion is numerically true as readily verified using the `prod2COP` function:

```
u <- c(0.41, 0.87); v <- c(0.13, 0.35); A <- -0.532; B <- 0.235
FGMcop( u,v, para= A*B / 3)
# 0.0521598638574___ 0.3034277347150___
prod2COP(u,v, cop1=FGMcop, para1=A, cop2=FGMcop, para2=B)
# 0.0521598638312605 0.3034277344807909
```

Author(s)

W.H. Asquith

References

Darsow, W.F., Nguyen, B., and Olsen, E.T., 1992, Copulas and Markov processes: Illinois Journal of Mathematics, v. 26, pp. 600–624, [doi:10.1215/IJM/1255987328](https://doi.org/10.1215/IJM/1255987328).

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[COP](#), [composite1COP](#), [composite2COP](#), [composite3COP](#), [convexCOP](#), [convex2COP](#), [glueCOP](#)

Examples

```
## Not run:
# Product P * N4212 ---> P (by identity)
u <- c(0.41, 0.87); v <- c(0.13, 0.35)
prod2COP(u,v, cop1=P, cop2=N4212cop, para1=NA, para2=2.12) # 0.0533 and 0.3045
COP(u,v, cop=P) # 0.0533 and 0.3045
## End(Not run)

## Not run:
para <- list(cop1=PLcop, para1=0.19, cop2=PLcop, para2=34.5)
UV <- simCOP(n=1000, cop=prod2COP, para=para, resamv01=FALSE, showresamv01=FALSE)
# This is large simulation run (with a lot of numerical operations) is expected
# at least for the Placketts and chosen parameters to trigger one or more NAs
# from derCOPinv(). The simCOP() function simply continues on with ignoring the
# solution or lack thereof for certain combinations, and simCOP() will report how
# many of the simulated values for sample of size n were computed. For example,
# for one n=1000, some 965 simulated values were returned. The defaults require
# that NAs, empty simulations, remain intact. We can try resampling:
UV <- simCOP(n=1000, cop=prod2COP, para=para, resamv01=TRUE, showresamv01=TRUE)
rhoCOP(cop=prod2COP, para=para) # -0.4271195 (theoretical)
rhoCOP(para=UV, as.sample=TRUE) # -0.4274703 #
## End(Not run)

## Not run:
para <- list(cop1=PLcop, para1=0.19, cop2=PLcop, para2=34.5)
# The prod2COP() might be one of the more sensitive to NAs in simulation because
# of the two partial numerical derivatives involved.
para$pinterval <- c(0.4, 0.6) # totally inappropriate interval for the integral
# for the prod2COP() definition. Because the ... are used so extensively, we have
# the "pinterval" for this function so that interval itself can be passed also.
UV <- simCOP(n=1000, cop=prod2COP, para=para, resamv01=TRUE, showresamv01=TRUE,
  pinterval=c(0, 1 ))
UV <- simCOP(n=1000, cop=prod2COP, para=para, resamv01=TRUE, showresamv01=TRUE,
  pinterval=c(0.4, 0.6)) #
## End(Not run)
```

Description

Kiriliouk *et al.* (2016, pp. 358–360) describe a *pseudo-polar* representation of bivariate data as a means to explore right-tail extremal dependency between the variables. Let (X_i, Y_i) (real values) or (U_i, V_i) (as probabilities) for $i = 1, \dots, n$ be a bivariate sample of size n . When such data are transformed into a “unit-Pareto” scale by

$$\hat{X}_i^* = n/(n+1 - R_{X,i}) \text{ and } \hat{Y}_i^* = n/(n+1 - R_{Y,i}),$$

where R is `rank()`, then letting each *component sum* or *pseudo-polar radius* be defined as

$$\hat{S}_i = \hat{X}_i^* + \hat{Y}_i^*,$$

and each respective *pseudo-polar angle* be defined as

$$\hat{W}_i = \hat{X}_i^* / (\hat{X}_i^* + \hat{Y}_i^*) = \hat{X}_i^* / \hat{S}_i,$$

a pseudo-polar representation is available for study.

A scatter plot of $\hat{W}_i$ (horizontal) versus $\hat{S}_i$ (vertical) will depict a *pseudo-polar plot* of the data. Kiriliouk *et al.* (2016) approach the pseudo-polar concept as a means to study extremal dependency in the sense of what are the contributions of the X and Y to their sum conditional on the sum being large. The *largeness* of $\hat{S}_i$ is assessed by its empirical cumulative distribution function and a threshold S_f stemming from f as a nonexceedance probability $f \in [0, 1]$.

A density plot of the $\hat{W}_i$ is a representation of extremal dependence. If the density plot shows low density for pseudo-polar angles away from 0 and 1 or bimodality on the edges then weak extremal dependency is present. If the density is substantial and uniform away from the the angles 0 and 1 or if the density peaks near $\hat{W} \approx 0.5$ then extremal dependency is strong.

Usage

```
psepolar(u, v=NULL, f=0.90, ...)
```

Arguments

u	Nonexceedance probability u in the X direction (actually the ranks are used so this can be a real-value argument as well);
v	Nonexceedance probability v in the Y direction (actually the ranks are used so this can be a real-value argument as well) and if NULL then u is treated as a two column R data.frame;
f	The nonexceedance probability of the distal $\hat{S}$ to flag in <code>Shat_ge_Sf</code> column of the output; and
...	Additional arguments to pass to the <code>dat2bernqua()</code> function of the lmomco package.

Value

An R data.frame is returned in the table element and the S_f is in the Sf element.

U	An echo of the u input;
V	An echo of the v input;
Xstar	The $\widehat{X}_i^*$ (Kiriliouk <i>et al.</i> , 2016, eq. 17.8, p. 359);
Ystar	The $\widehat{Y}_i^*$ (Kiriliouk <i>et al.</i> , 2016, eq. 17.8, p. 359);
FXhat1	The $F_{X,i} = 1 - 1/X_i^*$, which is the inverse of Kiriliouk <i>et al.</i> (2016, eq. 17.1, p. 354);
FYhat1	The $F_{Y,i} = 1 - 1/Y_i^*$, which is the inverse of Kiriliouk <i>et al.</i> (2016, eq. 17.1, p. 354);
FXhat3	The $F_{3,X,i} = (R_{X,i} - 0.5)/n$ corresponding to the “3” alternative identified by Kiriliouk <i>et al.</i> (2016, p. 365);
FYhat3	The $F_{3,Y,i} = (R_{Y,i} - 0.5)/n$ corresponding to the “3” alternative identified by Kiriliouk <i>et al.</i> (2016, p. 365);
What	The $\widehat{W}_i$ (Kiriliouk <i>et al.</i> , 2016, eq. 17.9, p. 359);
Shat	The $\widehat{S}_i$ (Kiriliouk <i>et al.</i> , 2016, eq. 17.9, p. 359); and
Shat_ge_Sf	A logical on whether the $\widehat{S}_i$ are larger than S_f .

Note

The default of $f=0.90$ means that the upper 90th percentile of the component sum will be identified in the output. This percentile is computed by the Bernstein empirical distribution function provided by the **lmomco** package through the `dat2bernqua()` function. Suggested arguments for ... are `poly.type="Bernstein"` and `bound.type="Carv"` though the former is redundant because it is the default of `dat2bernqua()`.

Author(s)

William Asquith <william.asquith@ttu.edu>

References

Kiriliouk, Anna, Segers, Johan, Warchoř, Michał, 2016, Nonparameteric estimation of extremal dependence: *in* Extreme Value Modeling and Risk Analysis, D.K. Dey and Jun Yan *eds.*, Boca Raton, FL, CRC Press, ISBN 978-1-4987-0129-7.

See Also

[spectralmeas](#), [stabtaildepf](#)

Examples

```
## Not run:
pse <- psepolar(simCOP(n=799, cop=PARETOcop, para=4.3,graphics=FALSE),bound.type="Carv")
pse <- pse$table # The Pareto copula has right-tail extreme dependency
plot(1/(1-pse$U), 1/(1-pse$V), col=pse$Shat_ge_Sf+1, lwd=0.8, cex=0.5, log="xy", pch=16)
plot(pse$What, pse$Shat, log="y", col=pse$Shat_ge_Sf+1, lwd=0.8, cex=0.5, pch=16)
plot(density(pse$What[pse$Shat_ge_Sf]), pch=16, xlim=c(0,1)) # then try the
# non-right tail extremal copula PSP as cop=PSP in the above psepolar() call.
## End(Not run)
```

PSP

The Ratio of the Product Copula to Summation minus Product Copula

Description

Compute *PSP copula* (Nelsen, 2006, p. 23) is named by the author (Asquith) for the **copBasic** package and is

$$\mathbf{PSP}(u, v) = \frac{\mathbf{\Pi}}{\mathbf{\Sigma} - \mathbf{\Pi}} = \frac{uv}{u + v - uv},$$

where $\mathbf{\Pi}$ is the *independence* or *product copula* (**P**) and $\mathbf{\Sigma}$ is the sum $\mathbf{\Sigma} = u + v$. The $\mathbf{PSP}(u, v)$ copula is a special case of the **N4212**(u, v) copula (**N4212cop**). The **PSP** is included in **copBasic** because of its simplicity and for pedagogical purposes. The name “PSP” comes from “Product, Summation, Product” to loosely reflect the mathematical formula shown. Nelsen (2006, p. 114) notes that the PSP copula shows up in several families and designates it as “ $\mathbf{\Pi}/(\mathbf{\Sigma} - \mathbf{\Pi})$.” The PSP is undefined for $u = v = 0$ but no internal trapping is made; calling functions will have to intercept the NaN so produced for $\{0, 0\}$. The **PSP** is left internally to not trap NaN so as to be available to stress other copula utility functions within the **copBasic** package.

Usage

```
PSP(u, v, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction; and
<code>...</code>	Additional arguments to pass, which for this copula are not needed, but given here to support flexible implementation.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[P](#), [N4212cop](#)

Examples

```
PSP(0.4, 0.6)
PSP(0, 0)
PSP(1, 1)
```

qua.regressCOP	<i>Perform Quantile Regression using a Copula by Numerical Derivative Method for V with respect to U</i>
----------------	--

Description

Perform *quantile regression* (Nelsen, 2006, pp. 217–218) using a copula by numerical derivatives of the copula ([derCOPinv](#)). If X and Y are random variables having quantile functions $x(F)$ and $y(G)$ and letting $y = \tilde{y}(x)$ denote a solution to $\Pr[Y \leq y \mid X = x] = F$, where F is a nonexceedance probability. Then the curve $y = \tilde{y}(x)$ is the quantile regression curve of V or Y with respect to U or X , respectively. If $F = 1/2$, then *median regression* is performed ([med.regressCOP](#)). Using copulas, the quantile regression is expressed as

$$\Pr[Y \leq y \mid X = x] = \Pr[V \leq G(y) \mid U = F(x)] = \Pr[V \leq v \mid U = v] = \frac{\delta \mathbf{C}(u, v)}{\delta u},$$

where $v = G(y)$ and $u = F(x)$. The general algorithm is

1. Set $\delta \mathbf{C}(u, v) / \delta u = F$,
2. Solve the regression curve $v = \tilde{v}(u)$ (provided by [derCOPinv](#)), and
3. Replace u by $x(u)$ and v by $y(v)$.

The last step is optional as step two produces the regression in probability space, which might be desired, and step 3 actually transforms the probability regressions into the quantiles of the respective random variables.

Usage

```
qua.regressCOP(f=0.5, u=seq(0.01, 0.99, by=0.01), cop=NULL, para=NULL, ...)
```

Arguments

<code>f</code>	A single value of nonexceedance probability F to perform regression at and defaults to median regression $F = 1/2$;
<code>u</code>	Nonexceedance probability u in the X direction;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula; and
<code>...</code>	Additional arguments to pass.

Value

An R data.frame of the regressed probabilities of V and provided $U = u$ values is returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[med.regressCOP](#), [derCOPinv](#), [qua.regressCOP.draw](#)

Examples

```
## Not run:
# Use a positively associated Plackett copula and perform quantile regression
theta <- 10
R <- qua.regressCOP(cop=PLACKETTcop, para=theta) # 50th percentile regression

plot(R$U,R$V, type="l", lwd=6, xlim=c(0,1), ylim=c(0,1), col=8)
lines(R$U,(1+(theta-1)*R$U)/(theta+1), col=4, lwd=1) # theoretical for Plackett, see
# (Nelsen, 2006, p. 218)
R <- qua.regressCOP(f=0.90, cop=PLACKETTcop, para=theta) # 90th-percentile regression
lines(R$U,R$V, col=2, lwd=2)
R <- qua.regressCOP(f=0.10, cop=PLACKETTcop, para=theta) # 10th-percentile regression
lines(R$U,R$V, col=3, lty=2)
mtext("Quantile Regression V wrt U for Plackett copula")#
## End(Not run)

## Not run:
# Use a composite copula with two Placketts with compositing parameters alpha and beta.
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
             para1=0.04, para2=5, alpha=0.9, beta=0.6)
plot(c(0,1),c(0,1), type="n", lwd=3,
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY")
# Draw the regression of V on U and then U on V (wrtV=TRUE)
qua.regressCOP.draw(cop=composite2COP, para=para, ploton=FALSE)
qua.regressCOP.draw(cop=composite2COP, para=para, wrtV=TRUE, lty=2, ploton=FALSE)
```

```

mtext("Composition of Two Plackett Copulas and Quantile Regression")#
## End(Not run)

## Not run:
# Use a composite copula with two Placketts with compositing parameters alpha and beta.
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
             para1=0.34, para2=50, alpha=0.63, beta=0.47)
D <- simCOP(n=3000, cop=composite2COP, para=para, cex=0.5)
qua.regressCOP.draw(cop=composite2COP, para=para, ploton=FALSE)
qua.regressCOP.draw(cop=composite2COP, para=para, wrtV=TRUE, lty=2, ploton=FALSE)
level.curvesCOP(cop=composite2COP, para=para, ploton=FALSE)
mtext("Composition of Two Plackett Copulas, Level Curves, Quantile Regression")

para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, # Note the singularity
             para1=0, para2=500, alpha=0.63, beta=0.47)
D <- simCOP(n=3000, cop=composite2COP, para=para, cex=0.5)
qua.regressCOP.draw(cop=composite2COP, para=para, ploton=FALSE)
qua.regressCOP.draw(cop=composite2COP, para=para, wrtV=TRUE, lty=2, ploton=FALSE)
level.curvesCOP(cop=composite2COP, para=para, ploton=FALSE)
mtext("Composition of Two Plackett Copulas, Level Curves, Quantile Regression")

pdf("quantile_regression_test.pdf")
for(i in 1:10) {
  para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, alpha=runif(1), beta=runif(1),
               para1=10^runif(1,min=-4,max=0), para2=10^runif(1,min=0,max=4))
  txts <- c("Alpha=", round(para$alpha, digits=4),
            "; Beta=", round(para$beta, digits=4),
            "; Theta1=", round(para$para1[1], digits=5),
            "; Theta2=", round(para$para2[1], digits=2))

  D <- simCOP(n=3000, cop=composite2COP, para=para, cex=0.5, col=3)
  mtext(paste(txts, collapse=""))
  qua.regressCOP.draw(f=c(seq(0.05, 0.95, by=0.05)),
                     cop=composite2COP, para=para, ploton=FALSE)
  qua.regressCOP.draw(f=c(seq(0.05, 0.95, by=0.05)),
                     cop=composite2COP, para=para, wrtV=TRUE, ploton=FALSE)
  level.curvesCOP(cop=composite2COP, para=para, ploton=FALSE)
}
dev.off() # done
## End(Not run)

```

qua.regressCOP.draw	<i>Draw Quantile Regressions using a Copula by Numerical Derivative Method for V with respect to U or U with respect to V</i>
---------------------	---

Description

Draw a suite of lines for specified nonexceedance probabilities representing the *quantile regression* (Nelsen, 2006, pp. 217–218) of either V with respect to U or U with respect to V depending upon an argument setting.

Usage

```
qua.regressCOP.draw(f=seq(0.1, 0.9, by=0.1), fs=0.5, cop=NULL, para=NULL,
                    ploton=TRUE, wrtV=FALSE, col=c(4,2), lwd=c(1,2), lty=1,...)
```

Arguments

<code>f</code>	Nonexceedance probability F to perform quantile regression at and defaults to a 10-percent-interval sequence. This vectorization of <code>f</code> for this function differs from that in qua.regressCOP and qua.regressCOP2 ;
<code>fs</code>	A special value of nonexceedance probability to draw with second values to arguments <code>col</code> and <code>lwd</code> and defaults to the median ($F = 1/2$);
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>ploton</code>	A logical to toggle on the plot;
<code>wrtV</code>	If <code>wrtV=FALSE</code> call qua.regressCOP and perform quantile regression of V with respect to U and if <code>wrtV=TRUE</code> call qua.regressCOP2 and perform regression of U with respect to V ;
<code>col</code>	A vector of two values for the color of the line to draw, where the first value is used for the <code>f</code> probabilities and the second value is used for the <code>fs</code> probability;
<code>lwd</code>	A vector of two values for the line width of the line to draw, where the first value is used for the <code>f</code> probabilities and the second value is used for the <code>fs</code> probability;
<code>lty</code>	The line type to draw; and
<code>...</code>	Additional arguments to pass.

Value

No values are returned, this function is used for its side effects.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[qua.regressCOP](#), [qua.regressCOP2](#)

Examples

```
# See example in qua.regressCOP documentation
```

qua.regressCOP2	<i>Perform Quantile Regression using a Copula by Numerical Derivative Method for U with respect to V</i>
-----------------	--

Description

Perform *quantile regression* (Nelsen, 2006, pp. 217–218) using a copula by numerical derivatives of the copula ([derCOPinv2](#)). If X and Y are random variables having quantile functions $x(F)$ and $y(G)$ and letting $x = \tilde{x}(y)$ denote a solution to $\Pr[X \leq x \mid Y = y] = F$, where F is a nonexceedance probability. Then the curve $x = \tilde{x}(y)$ is the quantile regression curve of U or X with respect to V or Y , respectively. If $F = 1/2$, then *median regression* is performed ([med.regressCOP2](#)). Using copulas, the quantile regression is expressed as

$$\Pr[X \leq x \mid Y = y] = \Pr[U \leq F(x) \mid V = F] = \Pr[U \leq u \mid V = F] = \frac{\delta \mathbf{C}(u, v)}{\delta v},$$

where $v = G(y)$ and $u = F(x)$. The general algorithm is

1. Set $\delta \mathbf{C}(u, v) / \delta v = F$,
2. Solve the regression curve $u = \tilde{u}(v)$ (provided by [derCOPinv2](#)), and
3. Replace u by $x(u)$ and v by $y(v)$.

The last step is optional as step two produces the regression in probability space, which might be desired, and step 3 actually transforms the probability regressions into the quantiles of the respective random variables.

Usage

```
qua.regressCOP2(f=0.5, v=seq(0.01,0.99, by=0.01), cop=NULL, para=NULL, ...)
```

Arguments

f	A single value of nonexceedance probability F to perform regression at and defaults to median regression $F = 1/2$;
v	Nonexceedance probability v in the Y direction;
cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula; and
...	Additional arguments to pass.

Value

An R data.frame of the regressed probabilities of U and $V = v$ is returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[med.regressCOP2](#), [derCOPinv2](#)

Examples

```
## Not run:
# Use a positively associated Plackett copula and perform quantile regression
theta <- 0.10
R <- qua.regressCOP2(cop=PLACKETTcop, para=theta) # 50th percentile regression
plot(R$U,R$V, type="l", lwd=6, xlim=c(0,1), ylim=c(0,1), col=8)
lines((1+(theta-1)*R$V)/(theta+1),R$V, col=4, lwd=1) # theoretical for Plackett,
# compare the theoretical form to that in qua.regressCOP---just switch terms around
# because of symmetry
R <- qua.regressCOP2(f=0.90, cop=PLACKETTcop, para=theta) # 90th-percentile regression
lines(R$U,R$V, col=2, lwd=2)
R <- qua.regressCOP2(f=0.10, cop=PLACKETTcop, para=theta) # 10th-percentile regression
lines(R$U,R$V, col=2, lty=2)
mtext("Quantile Regression U wrt V for Plackett copula")#
## End(Not run)
```

RAYcop

The Rayleigh Copula

Description

The *Rayleigh copula* (Boškoskia and others, 2018) is

$$\begin{aligned} \mathbf{C}_\Theta(u, v) &= \mathbf{RAY}(u, v; \Theta) = 1 + A - B, \\ A &= e^{\Theta a_2 - a_2} \left(e^{-a_1} \int_0^{\Theta a_2} e^{-s} I_0(2\sqrt{a_1 s}) \, ds - 1 \right) ds, \\ B &= e^{-a_1} \int_0^{a_2} e^{-s} I_0(2\sqrt{\Theta a_1 s}) \, ds, \end{aligned}$$

where $a_1 = -\log(1 - u)/(1 - \Theta)$, $a_2 = -\log(1 - v)/(1 - \Theta)$, $I_\nu(x)$ is the modified Bessel function of the first kind of order ν (see `base::besselI()`), and $\Theta \in (0, 1]$. The copula, as $\Theta \rightarrow 0^+$, limits to the *independence coupla* ($\mathbf{\Pi}(u, v)$; [P](#)) and as $\Theta \rightarrow 1^-$ limits to the *comonotonicity copula* ($\mathbf{M}(u, v)$; [M](#)). Finally, there are formulations of the Rayleigh copula using the *Marcum-Q function*, but the **copBasic** developer has not been able to make such work. If the Marcum-Q function could be used, then only one integration and not the two involving the modified Bessel function are possible. Infinite integrations begin occurring in the upper right corner for about $\Theta > 0.995$ at which point the $\mathbf{M}(u, v)$ copula is called in the source code.

Usage

```
RAYcop(u, v, para=NULL, rho=NULL, method=c("default"),
       rel.tol=.Machine$double.eps^0.5, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula;
<code>rho</code>	Value for Spearman Rho from which parameter Θ is computed by polynomial approximation and returned. The estimation appears sufficient for most practical applications (see Examples);
<code>method</code>	The computational method of integrals associated with the definition of the copula; this is designed for the ability to switch eventually in sources to Marcum-Q function implementation. The definition in January 2023 and default is to call the two Bessel function integrals shown for the definition in this documentation;
<code>rel.tol</code>	Argument of the same name for <code>integrate()</code> call; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Note

Documentation in Zeng and others [Part II] appear to have corrected the Marcum-Q function solution to the copula. The essence of that solution is with a Chi-distribution computation the Marcum-Q. Testing indicates that they have the correct solution, but the derivative for the conditional simulation as built into the design of **copBasic** has difficulties. Perhaps this is related to numerical precision of the Marcum-Q?

```
sapply(seq_len(length((u))), function(i) {
  a1 <- -log(1-u[i])
  if(is.infinite(a1)) return(v[i])
  a2 <- -log(1-v[i])
  if(is.infinite(a2)) return(u[i])
  a1 <- exp(log(a1) - log(1-p))
  a2 <- exp(log(a2) - log(1-p))
  a3 <- marcumq.chi(sqrt(2* a1), sqrt(2*p*a2)) # Zeng and others (Part II)
  a4 <- marcumq.chi(sqrt(2*p*a1), sqrt(2* a2)) # Zeng and others (Part II)
  zz <- 1 + (1-v[i])*a3 - (1-u[i])*(1-a4)      # Zeng and others (Part II)
  zz[zz < 0] <- 0
  zz[zz > 1] <- 1
  return(zz)
})
```

Author(s)

W.H. Asquith

References

Boškoscia, P., Debenjaka, A., Boshkoskab, B.M., 2018, Rayleigh copula for describing impedance data with application to condition monitoring of proton exchange membrane fuel cells: *European Journal of Operational Research*, v. 266, pp. 269–277, doi:10.1016/j.ejor.2017.08.058.

Zeng, X., Ren, J., Wang, Z., Marshall, S., and Durrani, T., 2014, Copulas for statistical signal processing (Part I)—Extensions and generalization: *Signal Processing* v. 14, pp. 691–702, doi:10.1016/j.sigpro.2013.07.009.

Zeng, X., Ren, J., Sun, M., Marshall, S., and Durrani, T., 2014, Copulas for statistical signal processing (Part II)—Simulation, optimal selection and practical applications: *Signal Processing*, v. 94, pp. 681–690, doi:10.1016/j.sigpro.2013.07.006.

See Also

M, P

Examples

```
RAYcop(0.2, 0.8, para=0.8) # [1] 0.1994658 (by the dual Bessel functions)

RAYcop(0.8, 0.2, para=RAYcop(rho=rhoCOP(cop=RAYcop, para=0.8)))
# [1] 0.1994651 from polynomial conversion of Rho to Theta

## Not run:
# Recipe for assembling the Spearman Rho to Theta polynomial in sources.
Thetas <- seq(0, 0.999, by=0.001); RH0s <- NULL
for(p in Thetas) RH0s <- c(RH0s, rhoCOP(cop=RAYcop, para=p))
LM <- lm(Thetas ~ RH0s + I(RH0s^2) + I(RH0s^4) + I(RH0s^6) - 1 )
Rho2Theta <- function(rho) {
  coes <- c(1.32682824, -0.38876290, 0.09072305, -0.02921836)
  sapply(rho, function(r) coes[1]*r^1 + coes[2]*r^2 + coes[3]*r^4 + coes[4]*r^6)
}
plot(RH0s, Thetas, type="l", col=grey(0.8), lwd=12, lend=1,
     xlab="Spearman Rho", ylab="Rayleigh Copula Parameter Theta")
lines(RH0s, Rho2Theta(RH0s), col="red", lwd=2) #
## End(Not run)
```

Description

These data represent porosity and permeability data from laboratory analysis for Well-266 Reinecke Oil Field, Horseshoe Atoll, Texas as used for the outstanding article by Saller and Dickson (2011). Dr. A.H. Saller shared a CSV file with the author of the **copBasic** package sometime in 2011. These data are included in this package because of the instruction potential of the bivariate relation between the geologic properties of permeability and porosity.

Usage

```
data(ReineckeWell266)
```

Format

An R data.frame with

WELLNO The number of the well, no. 266;

DEPTH The depth in feet to the center of the incremental spacings of the data;

FRACDOLOMITE The fraction of the core sample that is dolomite, 0 is 100 percent limestone;

Kmax The maximum permeability without respect to orientation in millidarcies;

POROSITY The porosity of the core sample; and

DOLOMITE Is the interval treated as dolomite (1) or limestone (0).

References

Saller, A.H., Dickson, J.A., 2011, Partial dolomitization of a Pennsylvanian limestone buildup by hydrothermal fluids and its effect on reservoir quality and performance: AAPG Bulletin, v. 95, no. 10, pp. 1745–1762.

Examples

```
## Not run:
data(ReineckeWell266)
summary(ReineckeWell266) # show summary statistics
## End(Not run)
```

ReineckeWells

Porosity and Permeability Data for the Reinecke Oil Field, Horseshoe Atoll, Texas

Description

These data represent porosity and permeability data from laboratory analysis for the Reinecke Oil Field, Horseshoe Atoll, Texas as used for the outstanding article by Saller and Dickson (2011). Dr. A.H. Art Saller shared a CSV file with the author of the **copBasic** package sometime in 2011. These data are included in this package because of the instruction potential of the bivariate relation between the geologic properties of permeability and porosity.

Usage

```
data(ReineckeWells)
```

Format

An R data.frame with

DOLOMITE The fraction of the core sample that is dolomite, 0 is 100 percent limestone;

Kmax The maximum permeability without respect to orientation in millidarcies;

K90 The horizontal (with respect to 90 degrees of the borehole) permeability in millidarcies;

Kvert The vertical permeability in millidarcies; and

POROSITY The porosity of the core sample.

References

Saller, A.H., Dickson, J.A., 2011, Partial dolomitization of a Pennsylvanian limestone buildup by hydrothermal fluids and its effect on reservoir quality and performance: AAPG Bulletin, v. 95, no. 10, pp. 1745–1762.

Examples

```
## Not run:
data(ReineckeWells)
summary(ReineckeWells) # show summary statistics
## End(Not run)
```

RFcop

The Raftery Copula

Description

The *Raftery copula* (Nelsen, 2006, p. 172) is

$$C_{\Theta}(u, v) = \mathbf{RF}(u, v) = \mathbf{M}(u, v) + \frac{1 - \Theta}{1 + \Theta} \left(uv \right)^{1/(1-\Theta)} \left[1 - (\max\{u, v\})^{-(1+\Theta)/(1-\Theta)} \right],$$

where $\Theta \in (0, 1)$. The copula, as $\Theta \rightarrow 0^+$ limits, to the *independence coupla* ($\mathbf{P}(u, v)$; **P**), and as $\Theta \rightarrow 1^-$, limits to the *comonotonicity copula* ($\mathbf{M}(u, v)$; **M**). The parameter Θ is readily computed from *Spearman Rho* (**rhoCOP**) by $\rho_{\mathbf{C}} = \Theta(4 - 3\Theta)/(2 - \Theta)^2$ or from *Kendall Tau* (**tauCOP**) by $\tau_{\mathbf{C}} = 2\Theta/(3 - \Theta)$. However, this copula like others within the **copBasic** package can be reflected (rotated) at will with the **COP** abstraction layer to acquire negative or inverse dependency (*countermonotonicity*) (see the **Examples**).

Usage

```
RFcop(u, v, para=NULL, rho=NULL, tau=NULL, fit=c("rho", "tau"), ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula;
<code>rho</code>	Optional Spearman Rho from which the parameter will be estimated and presence of rho trumps tau;
<code>tau</code>	Optional Kendall Tau from which the parameter will be estimated;
<code>fit</code>	If <code>para</code> , <code>rho</code> , and <code>tau</code> are all NULL, then the <code>u</code> and <code>v</code> represent the sample. The measure of association by the <code>fit</code> declaration will be computed and the parameter estimated subsequently. The <code>fit</code> has no other utility than to trigger which measure of association is computed internally by the <code>cor</code> function in R; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned. Otherwise if either `rho` or `tau` is given, then the Θ is computed and a list having

<code>para</code>	The parameter Θ ;
<code>rho</code>	Spearman Rho if the <code>rho</code> is given; and
<code>tau</code>	Kendall Tau if the <code>tau</code> is given but also if both <code>rho</code> and <code>tau</code> are NULL as mentioned next.

and if `para=NULL` and `rho` and `tau=NULL`, then the values within `u` and `v` are used to compute Spearman Rho (`fit="rho"`) or Kendall Tau (`fit="tau"`) and then compute the parameter, and this is returned in the aforementioned list.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[M](#), [P](#)

Examples

```
# Lower tail dependency of Theta = 0.5 --> 2*(0.5)/(1+0.5) = 2/3 (Nelsen, 2006, p. 214)
taildepCOP(cop=RFcop, para=0.5)$lambdaL # 0.66667

## Not run:
# Simulate for a Spearman Rho of 0.7, then extract estimated Theta that internally
# is based on Kendall Tau of U and V, then convert estimate to equivalent Rho.
set.seed(1)
```

```

UV <- simCOP(1000, cop=RFcop, RFcop(rho=0.7)$para)
Theta <- RFcop(UV$U, UV$V, fit="tau")$para # 0.607544
Rho <- Theta*(4-3*Theta)/(2-Theta)^2 # 0.682255 (nearly 0.7) #
## End(Not run)

## Not run:
set.seed(1)
UV <- simCOP(1000, cop=COP, para=list(cop=RFcop, para=RFcop(rho=0.5)$para, reflect=3))
cor(UV$U, UV$V, method="spearman") # -0.492677 as expected with reversal of V #
## End(Not run)

```

rhobevCOP

A Dependence Measure for a Bivariate Extreme Value Copula based on the Expectation of the Product of Negated Log-Transformed Random Variables U and V

Description

Compute a dependence measure based on the expectation of the product of transformed random variables U and V , which unnamed by Joe (2014, pp. 383–384) but symbolically is ρ_E , having a *bivariate extreme value copula* $\mathbf{C}_{BEV}(u, v)$ by

$$\rho_E = E[(-\log U) \times (-\log V)] - 1 = \int_0^1 [B(w)]^{-2} dw - 1,$$

where $B(w) = A(w, 1 - w)$, $B(0) = B(1) = 1$, $B(w) \geq 1/2$, and $0 \leq w \leq 1$, and where only bivariate extreme value copulas can be written as

$$\mathbf{C}_{BEV}(u, v) = \exp[-A(-\log u, -\log v)],$$

and thus in terms of the coupla

$$B(w) = -\log[\mathbf{C}_{BEV}(\exp[-w], \exp[w - 1])].$$

Joe (2014, p. 383) states that ρ_E is the correlation of the “survival function of a bivariate min-stable exponential distribution,” which can be assembled as a function of $B(w)$. Joe (2014, p. 383) also shows the following expression for *Spearman Rho*

$$\rho_S = 12 \int_0^1 [1 + B(w)]^{-2} dw - 3,$$

in terms of $B(w)$. This expression, in conjunction with [rhoCOP](#), was used to confirm the prior expression shown here for $B(w)$ in terms of $\mathbf{C}_{BEV}(u, v)$. Lastly, for *independence* ($uv = \mathbf{\Pi}$; [P](#)), $\rho_E = 0$ and for the *Fréchet–Hoeffding upper-bound copula* (perfect positive association), $\rho_E = 1$.

Usage

```
rhobevCOP(cop=NULL, para=NULL, as.sample=FALSE, brute=FALSE, delta=0.002, ...)
```

Arguments

cop	A bivariate extreme value copula function—the function rhobevCOP makes no provision for verifying whether the copula in cop is actually an <i>extreme value copula</i> ;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
as.sample	A logical controlling whether an optional R data.frame in para is used to compute a $\hat{\rho}_E$ by mean() of the product of negated log()'s in R. The user is required to cast para into estimated probabilities (see Examples);
brute	Should brute force be used instead of two nested integrate() functions in R to perform the double integration;
delta	The dw for the brute force (brute=TRUE) integration; and
...	Additional arguments to pass.

Value

The value for ρ_E is returned.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[rhoCOP](#), [tauCOP](#)

Examples

```
Theta <- GHcop(tau=1/3)$para      # Gumbel-Hougaard copula with Kendall Tau = 1/3
rhobevCOP(cop=GHcop, para=Theta) # 0.3689268 (RhoE after Joe [2014])
rhoCOP(  cop=GHcop, para=Theta) # 0.4766613 (Spearman Rho)

## Not run:
set.seed(394)
Theta <- GHcop(tau=1/3)$para      # Gumbel-Hougaard copula with Kendall Tau = 1/3
simUV <- simCOP(n=30000, cop=GHcop, para=Theta, graphics=FALSE) # large simulation
samUV <- simUV * 150; n <- length(samUV[,1]) # convert to fake unit system
samUV[,1] <- rank(simUV[,1]-0.5)/n; samUV[,2] <- rank(simUV[,2]-0.5)/n # hazen
rhobevCOP(para=samUV, as.sample=TRUE) # 0.3708275
## End(Not run)
```

Description

Compute the measure of association known as the *Spearman Rho* $\rho_{\mathbf{C}}$ of a copula according to Nelsen (2006, pp. 167–170, 189, 208) by

$$\rho_{\mathbf{C}} = 12 \int \int_{\mathcal{I}^2} \mathbf{C}(u, v) \, du dv - 3,$$

or

$$\rho_{\mathbf{C}} = 12 \int \int_{\mathcal{I}^2} [\mathbf{C}(u, v) - uv] \, du dv,$$

where the later equation is implemented by rhoCOP as the default method (method="default"). This equation, here having $p = 1$ and $k_p(1) = 12$, is generalized under [hoefCOP](#). The absence of the 12 in the above equation makes it equal to the *covariance of a copula* defined by the *Hoeffding Identity* (Joe, 2014, p. 54):

$$\text{cov}(U, V) = \int \int_{\mathcal{I}^2} [\mathbf{C}(u, v) - uv] \, du dv \text{ or}$$

$$\text{cov}(U, V) = \int \int_{\mathcal{I}^2} [\hat{\mathbf{C}}(u, v) - uv] \, du dv, \text{ which is}$$

$$\text{cov}(U, V) = \int \int_{\mathcal{I}^2} [u + v - 1 + \mathbf{C}(1 - u, 1 - v) - uv] \, du dv.$$

Depending on copula family (Joe, 2014, pp. 56 and 267), the alternative formulation for $\rho_{\mathbf{C}}$ could be used

$$\rho_{\mathbf{C}} = 3 - 12 \int \int_{\mathcal{I}^2} u \frac{\delta \mathbf{C}(u, v)}{\delta u} \, du dv = 3 - 12 \int \int_{\mathcal{I}^2} v \frac{\delta \mathbf{C}(u, v)}{\delta v} \, du dv,$$

where the first integral form corresponds to Joe (2014, eq. 248, p. 56) and is the method="joe21", and the second integral form is the method="joe12".

The integral

$$\int \int_{\mathcal{I}^2} \mathbf{C}(u, v) \, du dv,$$

represents the “volume under the graph of the copula and over the unit square” (Nelsen, 2006, p. 170) and therefore $\rho_{\mathbf{C}}$ is simple a rescaled volume under the copula. The second equation for $\rho_{\mathbf{C}}$ expresses the “average distance” between the joint distribution and statistical *independence* $\Pi = uv$. Nelsen (2006, pp. 175–176) shows that the following relation between $\rho_{\mathbf{C}}$ and $\tau_{\mathbf{C}}$ ([tauCOP](#)) exists

$$-1 \leq 3\tau - 2\rho \leq 1.$$

Usage

```
rhoCOP(cop=NULL, para=NULL, method=c("default", "joe21", "joe12"),
       as.sample=FALSE, brute=FALSE, delta=0.002, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
method	The form of integration used to compute (see above);
as.sample	A logical controlling whether an optional R data.frame in para is used to compute the $\hat{\rho}$ by dispatch to cor() function in R with method = "spearman";
brute	Should brute force be used instead of two nested integrate() functions in R to perform the double integration;
delta	The du and dv for the brute force integration using brute; and
...	Additional arguments to pass.

Value

The value for ρ_C is returned.

Note

Technically, Nelsen (2006) also shows that these definitions are a form of call to a *concordance function* $Q(C_1, C_2)$ of two copulas that involve $C_1 = C(u, v)$ and $C_2 = \Pi$. As such in order to keep rhoCOP a small function when brute=TRUE, ρ_C is computed by a special call to [tauCOP](#), which by itself and although titled for computation of *Kendall Tau*, does support the concordance function $Q(C_1, C_2)$ [see Nelsen (2006, pp. 158–159)] when given two different copulas and respective parameters as arguments. The well-known *Pearson correlation coefficient* equals Spearman rho value if random variables X and Y are both uniformly distributed on $[0, 1]$.

Author(s)

W.H. Asquith

References

- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
 Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[blomCOP](#), [footCOP](#), [giniCOP](#), [hoefCOP](#), [tauCOP](#), [wolfCOP](#), [joeskewCOP](#), [uvmoms](#)

Examples

```
rhoCOP(cop=PSP)           # 0.4784176

## Not run:
  n <- 1E4; set.seed(4784)  # quick Monte Carlo integration
  12*mean(replicate(n/10, sum(PSP(runif(n),runif(n)))/n)) - 3 # 0.4779 (for instance)
## End(Not run)

## Not run:
```

```

rhoCOP(cop=PSP, brute=TRUE) # 0.4684063
# Heavy CPU example showing dual-integration (fast) results in Rho mimicking the
# sample version
do_rho <- function(n) {
  uv <- simCOP(n=n, cop=PSP, ploton=FALSE, points=FALSE)
  return(cor(uv$U, uv$V, method="spearman"))
}
set.seed(1)
rhos <- replicate(200, do_rho(1000))
rho_sample <- mean(rhos); print(rho_sample) # 0.4764914
## End(Not run)

## Not run:
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
             para1=0.00395, para2=4.67, alpha=0.9392, beta=0.5699)
rhoCOP(cop=composite2COP, para=para) # -0.5924796

para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
             para1=0.14147, para2=20.96, alpha=0.0411, beta=0.6873)
rhoCOP(cop=composite2COP, para=para) # 0.2818874

para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
             para1=0.10137, para2=4492.87, alpha=0.0063, beta=0.0167)
rhoCOP(cop=composite2COP, para=para) # 0.9812919
rhoCOP(cop=composite2COP, para=para, brute=TRUE) # 0.9752155
## End(Not run)

## Not run:
# This is the same composited copula used in a highly asymmetric multi-modal
# plotting example under densityCOPplot(). Let us use that copula as a means to
# check on the Spearman Rho from alternative formulations by Joe (2014).
para <- list(alpha=0.15, beta=0.90, kappa=0.06, gamma=0.96,
             cop1=GHcop, cop2=PLACKETTcop, para1=5.5, para2=0.07)
"rhoCOPbyJoe21" <- function(cop=NULL, para=NULL, ...) { # Joe (2014, eq. 2.48)
  myint <- NULL
  try(myint <- integrate(function(u) {
    sapply(u,function(u) { integrate(function(v) {
      u * derCOP( u, v, cop=cop, para=para, ...)}, 0, 1)$value })), 0, 1))
  ifelse(is.null(myint), return(NA), return(3 - 12*myint$value))
}
"rhoCOPbyJoe12" <- function(cop=NULL, para=NULL, ...) { # Not in Joe (2014)
  myint <- NULL
  try(myint <- integrate(function(u) {
    sapply(u,function(u) { integrate(function(v) {
      v * derCOP2( u, v, cop=cop, para=para, ...)}, 0, 1)$value })), 0, 1))
  ifelse(is.null(myint), return(NA), return(3 - 12*myint$value))
}
rhoCOP(      cop=composite2COP, para=para) # 0.1031758
rhoCOPbyJoe21(cop=composite2COP, para=para) # 0.1031803
rhoCOPbyJoe12(cop=composite2COP, para=para) # 0.1031532
## End(Not run)

```

rmseCOP

Root Mean Square Error between a Fitted Copula and an Empirical Copula

Description

Compute the *root mean square error* RMSE_C (Chen and Guo, 2019, p. 29), which is computed using *mean square error* MSE as

$$\text{MSE}_C = \frac{1}{n} \sum_{i=1}^n (\mathbf{C}_n(u_i, v_i) - \mathbf{C}_{\Theta_m}(u_i, v_i))^2 \text{ and}$$

$$\text{RMSE}_C = \sqrt{\text{MSE}_C},$$

where $\mathbf{C}_n(u_i, v_i)$ is the *empirical copula* (empirical joint probability) for the i th observation, $\mathbf{C}_{\Theta_m}(u_i, v_i)$ is the fitted copula having m parameters in Θ . The $\mathbf{C}_n(u_i, v_i)$ comes from [EMPIRCop](#). The RMSE_C is in effect saying that the best copula will have its joint probabilities plotting on a 1:1 line with the empirical joint probabilities, which is an $\text{RMSE}_C = 0$. From the MSE_C shown above, the *Akaike information criterion* (AIC) [aicCOP](#) and *Bayesian information criterion* (BIC) [bicCOP](#) can be computed, which add a penalty for m parameters. These goodness-of-fits can assist in deciding one copula favorability over another, and another goodness-of-fit using the absolute differences between $\mathbf{C}_n(u, v)$ and $\mathbf{C}_{\Theta_m}(u, v)$ is found under [statTn](#).

Usage

```
rmseCOP(u, v=NULL, cop=NULL, para=NULL, ...)
```

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction; If not given, then a second column from argument u is attempted;
cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula; and
...	Additional arguments to pass to either copula (likely most commonly to the empirical copula).

Value

The value for RMSE_C is returned.

Author(s)

W.H. Asquith

References

Chen, Lu, and Guo, Shenglian, 2019, Copulas and its application in hydrology and water resources: Springer Nature, Singapore, ISBN 978–981–13–0574–0.

See Also

[EMPIRCop](#), [aicCOP](#), [bicCOP](#)

Examples

```
## Not run:
S <- simCOP(80, cop=GHcop, para=5) # Simulate some probabilities, but we
# must then treat these as data and recompute empirical probabilities.
U <- lmomco::pp(S$U, sort=FALSE); V <- lmomco::pp(S$V, sort=FALSE)
# The parent distribution is Gumbel-Hougaard extreme value copula.
# But in practical application we do not know that but say we speculate that
# perhaps the Galambos extreme value might be the parent. Then maximum
# likelihood is used to fit the single parameter.
pGL <- mleCOP(U,V, cop=GLcop, interval=c(0,20))$par

rmsees <- c(rmseCOP(U,V, cop=GLcop, para=pGL),
            rmseCOP(U,V, cop=P),
            rmseCOP(U,V, cop=PSP))
names(rmsees) <- c("GLcop", "P", "PSP")
print(rmsees) # We will see that the first RMSE is the smallest as the
# Galambos has the nearest overall behavior than the P and PSP copulas.
## End(Not run)
```

rN4220cop

The (Reflected) Copula of Equation 4.2.20 of Nelsen's Book

Description

The *rN4220 copula* (Nelsen, 2006, p. 118; eq. 4.2.20) is named by the author (Asquith) for the **copBasic** package and is a reflected version of [N4220cop](#) defined as

$$C_{rN4220}(u, v; \Theta) = u + v - 1 + \left[\log \left(\exp((1-u)^{-\Theta}) + \exp((1-v)^{-\Theta}) - \exp(1) \right) \right]^{-1/\Theta}.$$

The $rN4220(u, v)$ copula is *not comprehensive* because for $\Theta = 0^+$ the copula becomes the *independence* $P(u, v)$ copula ([P](#)) and as $\Theta \rightarrow \infty$ the copula becomes $M(u, v)$ ([M](#)). The copula is undefined for $\Theta < 0$. This copula has *upper-tail dependency*, which is the reverse of the $N4220(u, v)$ that has *lower-tail dependency*.

Usage

```
rN4220cop(u, v, para=NULL, insertM=TRUE, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula;
<code>insertM</code>	For the tail that pinches out towards $\mathbf{M}(u, v)$ (M) use an <i>ad hoc</i> algorithm to insert that copula into the tail (refer to source code) and that algorithm is dependent on the simCOP and the numerical partial derivative inversion for simulation that the package is based around; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

Examples

```
sapply(c(0.1, 1, 10), function(p) rN4220cop(0.4, 0.6, para=p))
# 0.2611771 0.3714941 0.4000000 # perfect correlation controls for p=10
sapply(c(0.1, 1, 10), function(p) rN4220cop(0.2, 0.1, para=p))
# 0.02331165 0.04762172 0.09998847
sapply(c(0.1, 1, 10), function(p) N4220cop(0.2, 0.1, para=p))
# 0.03549625 0.09993412 0.10000000
```

sectionCOP

The Sections or Derivative of the Sections of a Copula

Description

Compute the *copula sections* or the (partial) *derivatives of copula sections* of a copula (Nelsen, 2006, pp. 12–14). The *horizontal section* at $V = a$ (a constant) is

$$t \mapsto \mathbf{C}(t, a), \text{ and}$$

the *vertical section* at $U = a$ (a constant, with respect to V or `wrtV=TRUE`) is

$$t \mapsto \mathbf{C}(a, t).$$

The partial derivatives of the copula sections are *conditional cumulative distribution functions* (see [derCOP](#) and [derCOP2](#)). The derivatives are constrained as

$$0 \leq \frac{\delta}{\delta u} \mathbf{C}(u, v) \leq 1, \text{ and}$$

$$0 \leq \frac{\delta}{\delta v} \mathbf{C}(u, v) \leq 1.$$

Usage

```
sectionCOP(f, cop=NULL, para=NULL, wrtV=FALSE, dercop=FALSE, delt=0.005,
           ploton=TRUE, lines=TRUE, xlab="NONEXCEEDANCE PROBABILITY", ...)
```

Arguments

<code>f</code>	A single value of nonexceedance probability u or v along the horizontal U axis or vertical V axis of the unit square $\mathcal{I}^2$;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters, if needed, to pass to the copula;
<code>wrtV</code>	A logical to toggle between with respect to v or u (default). The default provides the vertical section whereas the horizontal comes from <code>wrtV = TRUE</code> ;
<code>dercop</code>	A logical that triggers the derivative of the section;
<code>delt</code>	The increment of the level curves to plot, defaults to 5-percent intervals;
<code>ploton</code>	A logical to toggle on the plot;
<code>lines</code>	Draw the lines of diagonal to the current device;
<code>xlab</code>	A label for the x-axis title passed to <code>plot()</code> in R; and
<code>...</code>	Additional arguments to pass to the <code>plot()</code> and <code>lines()</code> functions in R.

Value

An R list is returned.

<code>t</code>	The nonexceedance probability along the section. The nomenclature t mimics Nelsen (2006) and is <i>not</i> the same as the u or v ;
<code>seccop</code>	The section of the copula or its derivative;
<code>wrt</code>	A text string declaring what the setting for <code>wrtV</code> was;
<code>fvalue</code>	The provided value of nonexceedance probability; and
<code>isderivative</code>	A logical stating whether the derivative of the section is <code>seccop</code> .

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[COP](#), [diagCOP](#)

Examples

```
## Not run:
# EXAMPLE 1, plot the v=0.55 section and then u=0.55 section, which will overlay
# the other because the PSP is a symmetrical copula
tmp <- sectionCOP(0.55, cop=PSP, ylab="COPULA SECTIONS", lwd=5, col=2)
tmp <- sectionCOP(0.55, cop=PSP, wrtV=TRUE, ploton=FALSE, lwd=2, col=3)
# now add the v=0.85 section and the u=0.85, again overlay each other
tmp <- sectionCOP(0.85, cop=PSP, ploton=FALSE, lwd=5, col=2, lty=2)
tmp <- sectionCOP(0.85, cop=PSP, wrtV=TRUE, ploton=FALSE, lwd=2, col=3, lty=2)#
## End(Not run)

## Not run:
# EXAMPLE 2, v=0.35 section and derivative (the conditional distribution) function
tmp <- sectionCOP(0.35, cop=PSP, ylab="COPULA SECTIONS OR DERIV.", lwd=5, col=3)
tmp <- sectionCOP(0.35, cop=PSP, dercop=TRUE, ploton=FALSE, col=3)
# The thin green line represents the cumulative distribution function conditional
# on u = 0.35 from the derCOP function. Then see Example 3
## End(Not run)

## Not run:
# EXAMPLE 3 (random selection commented out)
#para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, alpha=runif(1), beta=runif(1),
#             para1=10^runif(1,min=-4, max=0), para2=10^runif(1,min= 0, max=4))
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, alpha=0.7, beta=0.22,
             para1=0.0155, para2=214.4)
txts <- c("Alpha=", round(para$alpha, digits=4),
          "; Beta=", round(para$beta, digits=4),
          "; Theta1=", round(para$para1[1], digits=5),
          "; Theta2=", round(para$para2[1], digits=2))
layout(matrix(1:2,byrow=TRUE))
D <- simCOP(n=1000, cop=composite2COP, para=para, cex=0.5, col=rgb(0,0,0,0.2), pch=16)
mtext(paste(txts,collapse=""))
#f <- c(runif(1),runif(1))
f <- c(0.2,0.9) # RED is the horizontal section and BLACK is the vertical section
segments(f[1],0,f[1],1, col=2, lwd=2); segments(0,f[2],1,f[2], lwd=2)
ftxt <- c("Sections (thick) and derivatives (thin) at ", f, " nonexceed. prob.")
tmp <- sectionCOP(f[1],cop=composite2COP,para=para, col=2, lwd=4)
tmp <- sectionCOP(f[1],cop=composite2COP,para=para, dercop=TRUE, ploton=FALSE, col=2)
tmp <- sectionCOP(f[2],cop=composite2COP,para=para,wrtV=TRUE,ploton=FALSE,lwd=4)
tmp <- sectionCOP(f[2],cop=composite2COP,para=para,wrtV=TRUE,ploton=FALSE,dercop=TRUE)
mtext(paste(ftxt, collapse=""))
# The thin lines are the CDFs conditional on the respective values of "f". Carefully
# compare the point densities along and near the sections in the top plot to the
# respective shapes of the CDFs in the bottom plot. If the bottom plot were rotated
# 90 degrees clockwise and then reflected top to bottom, the conditional quantile
# function QDF results. Reflection is needed because, by convention, QDFs are monotonic
# increasing to right---functions derCOPinv() and derCOPinv2() provide the CDF inversion.
## End(Not run)
```

Description

Compute the *lower semi-correlations* (bottom-left)

$$\rho_{\mathbf{C}}^{N--}(u, v; a) = \rho_N^{--}(a) \text{ and}$$

compute the *upper semi-correlations* (top-right)

$$\rho_{\mathbf{C}}^{N++}(u, v; a) = \rho_N^{++}(a)$$

of a copula $\mathbf{C}(u, v)$ (Joe, 2014, p. 73) using numerical simulation. The semi-correlations are defined as

$$\begin{aligned}\rho_N^{--}(a) &= \text{cor}[Z_1, Z_2 \mid Z_1 < -a, Z_2 < -a], \\ \rho_N^{++}(a) &= \text{cor}[Z_1, Z_2 \mid Z_1 > +a, Z_2 > +a], \text{ and} \\ \rho_N(a > -\infty) &= \text{cor}[Z_1, Z_2],\end{aligned}$$

where $\text{cor}[z_1, z_2]$ is the familiar *Pearson correlation function*, which is in **R** the syntax `cor(..., method="pearson")`, parameter $a \geq 0$ is a truncation point that identifies *truncated tail regions* (Joe, 2014, p. 73), and lastly $(Z_1, Z_2) \sim \mathbf{C}(\Phi, \Phi)$ and thus from the standard normal distribution $(Z_1, Z_2) = (\Phi^{-1}(u), \Phi^{-1}(v))$ where the random variables $(U, V) \sim \mathbf{C}$.

The semi-correlations are extended for the **copBasic** package into bottom right and top left versions as well by

$$\begin{aligned}\rho_N^{+-}(a) &= \text{cor}[Z_1, Z_2 \mid Z_1 > +a, Z_2 < -a], \text{ and} \\ \rho_N^{-+}(a) &= \text{cor}[Z_1, Z_2 \mid Z_1 < -a, Z_2 > +a].\end{aligned}$$

As a result, the notations $--$, $++$, $+-$, and $-+$ can be used to represent each of the respective corners bottom-left, top-right, bottom-right, and top-left of the (u, v) domain with the respective truncation. These words are used in the variable names of the returned list from the function.

Usage

```
semicorCOP(cop=NULL, para=NULL, truncation=0, n=0, as.sample=FALSE, ...)
```

Arguments

<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>truncation</code>	The truncation value for a , which is in standard normal variates, and the default of zero is the origin (medians);
<code>n</code>	The sample size n for simulation estimates of the ρ_N ;
<code>as.sample</code>	A logical controlling whether an optional <code>data.frame</code> in <code>para</code> is used to compute the $\hat{\rho}_N$ (see Note); and
<code>...</code>	Additional arguments to pass to the copula.

Value

The value(s) for ρ_N , ρ_N^{--} , ρ_N^{++} , ρ_N^{+-} , and ρ_N^{-+} are returned.

Note

The sample semi-correlations can be computed from a two-column table that is passed into the function using the `para` argument. Although the truncation point $a \geq 0$, as a increases and focus is increasingly made into one or the other truncated tail regions, the sample version with data becomes decreasing well estimated because the available sample size diminishes. The `para` argument can contain probabilities or raw data because internally the function computes the *Hazen plotting positions* (e.g. $u_i = (i - 0.5)/n$ for rank i and sample size n) because Joe (2014, pp. 9, 17, 245, 247–248) repeatedly emphasizes this form of plotting position when normal scores are involved.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[giniCOP](#), [rhoCOP](#), [tauCOP](#), [COP](#)

Examples

```
## Not run:
# Gumbel-Hougaard copula with Pearson rhoN = 0.4 (by definition)
run <- sapply(1:50, function(i) semicorCOP(cop=GHcop, para=1.350, n=600))
mean(unlist(run[1,])) # cor.normal.scores
mean(unlist(run[2,])) # botleft.semicor
mean(unlist(run[3,])) # topright.semicor
sd( unlist(run[1,])) # cor.normal.scores (These are our sampling variations
sd( unlist(run[2,])) # botleft.semicor for the n=600 used as a Monte
sd( unlist(run[3,])) # topright.semicor Carlo simulation.)
# The function returns: rhoN = 0.392112, rhoN--= 0.117674, rhoN+= 0.404733
# standard deviations (0.038883) (0.073392) (0.073942)
# Joe (2014, p. 72) shows: rhoN = 0.4____, rhoN--= 0.132___, rhoN+= 0.415___
# standard deviations (not avail) (0.08____) (0.07____)
# We see alignment with the results of Joe with his n=600. #
## End(Not run)

## Not run:
p <- 0.5 # Reasonable strong positive association for the Raftery copula,
# but then we are going to be reflecting and rotating the copula.
# See similar Example under COP() function.
"RFcop1" <- function(u,v, para) COP(u,v, cop=RFcop, para=para, reflect="1")
"RFcop2" <- function(u,v, para) COP(u,v, cop=RFcop, para=para, reflect="2")
"RFcop3" <- function(u,v, para) COP(u,v, cop=RFcop, para=para, reflect="3")
"RFcop4" <- function(u,v, para) COP(u,v, cop=RFcop, para=para, reflect="4")
RF <- NULL; n <- 10000
RF <- rbind(RF, as.data.frame(semicorCOP(cop=RFcop1, para=p, n=n))[1:5])
RF <- rbind(RF, as.data.frame(semicorCOP(cop=RFcop2, para=p, n=n))[1:5])
RF <- rbind(RF, as.data.frame(semicorCOP(cop=RFcop3, para=p, n=n))[1:5])
```

```

RF <- rbind(RF, as.data.frame(semicorCOP(cop=RFcop4, para=p, n=n))[,1:5])
print(RF[,1:3]) # total sample and lower and upper semi-correlations
# cor.normal.scores botleft.semicor topright.semicor
# 1      0.5587837      0.74124686      0.10027641
# 2      0.5567889      0.10302772      0.73729702
# 3     -0.5807201     -0.04683536     -0.01714573 # see near zeros --,++
# 4     -0.5698139      0.03040520      0.05125916 # see near zeros --,++
print(RF[,2:5]) # now look at all four corners
# botleft.semicor topright.semicor topleft.semicor botright.semicor
# 1      0.74124686      0.10027641      0.01529508      0.02046530
# 2      0.10302772      0.73729702     -0.05195628      0.01747874
# 3     -0.04683536     -0.01714573     -0.11106842     -0.74077321
# 4      0.03040520      0.05125916     -0.74516061     -0.07636233
# Notice how the tight tail of the copula, being reflected and rotated
# into each of the four corners, shows semi-correlation magnitude of 0.74.
# See the copula density plots of COP() Examples section.
## End(Not run)

```

simcomposite3COP	<i>Compute the L-comoments of a Four-Value Compositied Copula by Simulation</i>
------------------	---

Description

Simulate copula parameters and compute *L-comoments* and provision for plotting features for a composited copula using four compositing parameters (see [composite3COP](#)). The compositing parameters are each independent and uniformly distributed:

$$\alpha \sim U[0, 1]; \beta \sim U[0, 1]; \kappa \sim U[0, 1]; \gamma \sim U[0, 1].$$

L-comoment estimation is provided by the [lcomCOP](#).

Usage

```

simcomposite3COP(nsim=100, compositor=composite3COP,
                 parents=NULL, ploton=FALSE, points=FALSE,
                 showpar=FALSE, showresults=FALSE, digits=6, ...)

```

Arguments

nsim	Number of simulations to perform;
compositor	The compositing function that could be either composite1COP , composite2COP , and composite3COP ;
parents	A special parameter list (see Note);
ploton	A logical to toggle on intermediate plotting;
points	A logical to actually draw the simulations on the ploton by the points() function in R;
showpar	Print the simulated parameter set with each iteration;

showresults	Print the results (useful if harvest results from a batch operation in R);
digits	The number digits to pass to round if showresults is true; and
...	Additional arguments to pass.

Value

An R matrix of results is returned. Each row represents a single simulation run. The first four columns are the α , β , κ , and γ *compositing parameters* and are labeled as such. The next two columns are the opposing diagonals, by first row and then second, of the *L-comoment correlation*. The following two columns are the opposing diagonals, by row and then second, of the *L-coskew*. The following two columns are the opposing diagonals, by row and then second, of the *L-cokurtosis*. The L-comoment columns are labeled to reflect the L-comoment matrix: T2.21 means the L-comoment correlation row 2 column 1 and T3.12 mean the L-coskew row 1 column 2. The remaining columns represent the Θ_n parameters for copula 1, the Θ_m parameters for copula 2. The columns are labeled Cop1Thetas or Cop2Thetas.

Note

The following descriptions list in detail the parents argument structure and content of the para argument:

cop1 — Function of the first copula;
 cop2 — Function of the second copula;
 para1gen — Function to generate random parameters for the first copula; and
 para2gen — Function to generate random parameters for the second copula.

The para argument of this function are passed to the function contained in compositor and are therefore subject to further constraints in items should such constraints exist.

Author(s)

W.H. Asquith

References

Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.

See Also

[lcomCOP](#), [simcompositeCOP](#)

Examples

```
## Not run:
# EXAMPLE 1: Make a single simulation result.
mainpara <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
                 para1gen=function() { return(c(10^runif(1, min=-5, max=0))) },
                 para2gen=function() { return(c(10^runif(1, min= 0, max=5))) })
```

```

v <- simcompositeCOP(nsim=1, parent=mainpara, showresults=TRUE)
print(v)

# EXAMPLE 2: Make 1000 "results" and plot two columns.
mainpara <- list(cop1=PLACKETTcop, cop2=N4212cop,
  para1gen=function() { return(c(10^runif(1, min=-5, max=5))) },
  para2gen=function() { return(c(10^runif(1, min= 0, max=2))) })
v <- simcomposite3COP(nsim=100, parent=mainpara); labs <- colnames(v)
plot(v[,5], v[,7], # open circles are 1 with respect to 2
  xlab=paste(c(labs[5], "and", labs[6]), collapse=" "),
  ylab=paste(c(labs[6], "and", labs[8]), collapse=" "))
points(v[,6], v[,8], pch=16) # black dots are 2 with respect to 1
## End(Not run)

```

simcompositeCOP	<i>Compute the L-comoments of a Two-Value Composited Copula by Simulation</i>
-----------------	---

Description

Simulate copula parameters and compute *L-comoments* and provision for plotting features for a composited copula using using two compositing parameters (see [composite1COP](#) as well as [composite2COP](#)). The compositing parameters are each independent and uniformly distributed:

$$\alpha \sim U[0, 1]; \beta \sim U[0, 1].$$

L-comoment estimation is provided by the [lcomCOP](#).

Usage

```

simcompositeCOP(nsim=100, compositor=composite2COP,
  parents=NULL, ploton=FALSE, points=FALSE,
  showpar=FALSE, showresults=FALSE, digits=6, ...)

```

Arguments

nsim	Number of simulations to perform;
compositor	The compositing function, could be either composite1COP or composite2COP . Each of these is acceptable because two compositing parameters are used;
parents	A special parameter list (see Note);
ploton	A logical to toggle on intermediate plotting;
points	A logical to actually draw the simulations on the ploton by points() function in R;
showpar	Print the simulated parameter set with each iteration;
showresults	Print the results (useful if harvest results from a batch operation in R);
digits	The number digits to pass to round if showresults is true; and
...	Additional arguments to pass.

Value

An R matrix of results is returned. Each row represents a single simulation run. The first two columns are the α and β *compositing parameters* and are labeled as such. The next two columns are the opposing diagonals, by first row and then second, of the *L-comoment correlation*. The following two columns are the opposing diagonals, by row and then second, of the *L-coskew*. The following two columns are the opposing diagonals, by row and then second, of the *L-cokurtosis*. The L-comoment columns are labeled to reflect the L-comoment matrix: T2.21 means the L-comoment correlation row 2 column 1 and T3.12 mean the L-coskew row 1 column 2. The remaining columns represent the Θ_n parameters for copula 1, the Θ_m parameters for copula 2. The columns are labeled Cop1Thetas or Cop2Thetas.

Note

The following descriptions list in detail the parents argument structure and content of the para argument:

cop1 — Function of the first copula;

cop2 — Function of the second copula;

para1gen — Function to generate random parameters for the first copula; and

para2gen — Function to generate random parameters for the second copula.

The para argument of this function are passed to the function contained in compositor and are therefore subject to further constraints in items should such constraints exist.

Author(s)

W.H. Asquith

References

Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.

See Also

[lcomCOP](#), [simcomposite3COP](#)

Examples

```
## Not run:
# A single simulation result.
mainpara <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
                 para1gen=function() { return(c(10^runif(1,min=-5,max=0))) },
                 para2gen=function() { return(c(10^runif(1,min= 0,max=5))) })
v <- simcompositeCOP(nsim=1, parent=mainpara, showresults=TRUE)
print(v) # for review
## End(Not run)
```

simCOP

Simulate a Copula by Numerical Derivative Method

Description

Perform a simulation and visualization of a copula using numerical partial derivatives of the copula (Nelsen, 2006, p. 32). The method is more broadly known as *conditional simulation method*. Because a focus of **copBasic** is on copula theory for pedagogic purposes, the coupling between simulation and subsequent visualization is emphasized by this function by it providing for both simulation and plotting operations by default.

The `simCOP` function is based on a uniformly simulating nonexceedance probability u and then conditioning the v from the inverse of the sectional derivative for V with respect to U (see `derCOPinv`). The function for speed will only report a warning if at least one of the requested simulations in `n` could not be made because of `uniroot`'ing problems in `derCOPinv`. The returned `data.frame` will be shortened automatically, but this can be controlled by `na.rm`. Failure of a simulation is purely dependent failure of the derivative inversion. In general, inversion should be quite robust for continuous or near continuous copulas and even copulas with singularities should be more or less okay. Lastly, the logical combination `na.rm=FALSE` and `kept=TRUE` could be used to isolate those combinations giving `derCOPinv` problems. The implemented simulation method in the **copBasic** package is known as the *conditional distribution method* (Nelsen, 2006; pp. 40–41), *conditional method*, or *Rosenblatt transform* (Joe, 2014, p. 270).

Usage

```
simCOP(n=100, cop=NULL, para=NULL, na.rm=TRUE, seed=NULL, kept=FALSE,
       graphics=TRUE, ploton=TRUE, points=TRUE, snv=FALSE,
       infsnv.rm=TRUE, trapinfsnv=.Machine$double.eps,
       resamv01=FALSE, showresamv01=FALSE, ...)
rCOP(n, cop=NULL, para=NULL, na.rm=TRUE, seed=NULL,
     resamv01=FALSE, showresamv01=FALSE, ...)
```

Arguments

<code>n</code>	A sample size, default is $n = 100$;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters, if needed, to pass to the copula;
<code>na.rm</code>	A logical to toggle the removal of NA entries should they form on the returned <code>data.frame</code> . A well implemented copula should accommodate and not return NA but because this package relies on numerical derivation, it was decided to have a mechanism to handle this;
<code>seed</code>	The integer seed to pass immediately to <code>set.seed()</code> ;
<code>kept</code>	Keep the t uniform random variable for the simulation as the last column in the returned <code>data.frame</code> ;
<code>graphics</code>	A logical that will disable graphics by setting <code>ploton</code> and <code>points</code> to <code>FALSE</code> and overriding whatever their settings were;

ploton	A logical to toggle on the plot (see Examples in vuongCOP);
points	A logical to actually draw the simulations by the <code>points()</code> function in R;
snv	A logical to convert the $\{u, v\}$ to standard normal scores (variates) both for the optional graphics and the returned R <code>data.frame</code> . Curiously, Joe (2014) advocates extensively for use of normal scores, which is in contrast to Nelsen (2006) who does not;
infsnv.rm	A logical that will quietly strip out any occurrences of $u = \{0, 1\}$ or $v = \{0, 1\}$ from the simulations because these are infinity in magnitude when converted to standard normal variates has been selected. Thus, this logical only impacts logic flow when <code>snv</code> is TRUE. The <code>infsnv.rm</code> is mutually exclusive from <code>trapinfsnv</code> ;
trapinfsnv	If TRUE and presumably small, the numerical value of this argument (η) is used to replace $u = \{0, 1\}$ and $v = \{0, 1\}$ with $u(0) = v(0) = \eta$ or $u(1) = v(1) = 1 - \eta$ as appropriate when conversion to standard normal variates has been selected. The setting of <code>trapinfsnv</code> only is used if <code>snv</code> is TRUE and <code>infsnv.rm</code> is FALSE;
resamv01	A logical triggering resampling for the elements of $v = 0$ and $v = 1$ (see Examples). This is a relatively late addition to copBasic logic and hence is disabled by default. If this is set to true, then the operations related to <code>infsnv.rm</code> and <code>trapinfsnv</code> are never to be involved later down in the functions' logic;
showresamv01	A logical providing a trigger to display a <code>message()</code> within the resampling loops for $v \leq 0, v \geq 1$; and
...	Additional arguments to pass to the <code>points()</code> function in R.

Value

An R `data.frame` of the simulated values is returned.

Note

Function `rCOP` is a light-weight implementation for bivariate copula random variates that dispatches to [simCOPmicro](#) and bypasses the graphical and other features of `simCOP`. Finally, an experimental parallel for the empirical copula is [EMPIRsim](#). Note, the equivalent of a `resamv01` is not implemented at the level of [simCOPmicro](#) because it is better to do such at the abstraction level of `rCOP` and `simCOP`.

Author(s)

W.H. Asquith

References

- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
 Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[derCOPinv](#), [simCOPmicro](#)

Examples

```
simCOP(n=5, cop=PARETOcop, para=2.4)

# We can find some unusual simulation combinations for which V == 0 or 1 and might have
# downstream operations simply not able to handle infinite quantiles (say) at the edges.
# We can readily offload the issue back to the rCOP() or simCOP() with resamv01 argument.
nsim <- 800; para <- list(cop=PLcop, para=150, alpha=0.8, beta=0.3)
JK <- rCOP(nsim, cop=composite1COP, para=para, seed=1, resamv01=FALSE)
print(JK[JK[,2] == 1,]) # 189 0.9437248 1. So, row 189 in this example has V=1, and
UV <- rCOP(nsim, cop=composite1COP, para=para, seed=1, resamv01=TRUE ) # changing the
# resamv01 argument to TRUE, we get this message and no V=1 in the returned data frame.
# rCOP() has some v >= 1, resampling those <---- This is the message output.

## Not run:
# The simCOP function is oft used in other Examples sections through this package.
simCOP(n=10, cop=W) # Frechet lower-bound copula
simCOP(n=10, cop=P) # Independence copula
simCOP(n=10, cop=M, col=2) # Frechet upper-bound copula
simCOP(n=10, cop=PSP) # The PSP copula
## End(Not run)

## Not run:
# Now simulate the PSP copula, add the level curves of the copula, and demonstrate
# the uniform distribution of marginals on the correct axes (U [top] and V [left]).
D <- simCOP(n=400, cop=PSP) # store simulated values in D
level.curvesCOP(cop=PSP, ploton=FALSE)
rug(D$U, side=3, col=2); rug(D$V, side=4, col=2)

# Now let us get more complicated and mix two Plackett copulas together using the
# composite2COP as a "compositor." The parameter argument becomes more complex, but
# is passed as shown into composite2COP.
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, alpha=0.3, beta=0.5, para1=0.1, para2=50)
D <- simCOP(n=950, cop=composite2COP, para=para, col=grey(0, 0.2), pch=16, snv=TRUE) #
## End(Not run)
```

simCOPmicro

Simulate V from U through a Copula by Numerical Derivative Method

Description

Perform bivariate simulation of random but coupled variables V from U through a copula (Nelsen, 2006, p. 32) by inversion of the numerical derivatives of the copula ([derCOPinv](#), [derCOPinv2](#)). The method is more broadly known as *conditional simulation method*. An elaborate implementation is available in [simCOP](#), which unlike `simCOPmicro`, has provisions (default) for graphical support. The `simCOPmicro` function is intended to be a minimalist version for copula simulation, and such a version is useful for pedagogic purposes including *conditional distributions*, *conditional quantile functions*, and *copula reflection* (see **Note** and [COP](#)). An extended educational discussion of simulation using the conditional method is available in the **Note** section of [derCOPinv](#).

Some definitions are needed. The copula of $(1 - U, 1 - V)$ is the *survival copula* (**surCOP**) and is defined as

$$\hat{\mathbf{C}}(u, v) = u + v - 1 + \mathbf{C}(1 - u, 1 - v),$$

whereas, following the notation of Joe (2014, pp. 271–272), the copula of $(1 - U, V)$ is defined as

$$\dot{\mathbf{C}}(u, v) = v - \mathbf{C}(1 - u, v), \text{ and}$$

the copula of $(U, 1 - V)$ is defined as

$$\ddot{\mathbf{C}}(u, v) = u - \mathbf{C}(u, 1 - v).$$

Careful consideration of the nomenclature is necessary as confusion with the occurrences of $1 - u$ and $1 - v$ easily conflate meaning. The nomenclature for the *survival copula* is more elaborately shown under **surCOP**. The difficulty is that the bivariate arguments to the *survival copula* are *exceedance probabilities*.

For simulation, again following the nomenclature of Joe (2014, p. 272), the conditional distribution functions (numerical derivatives; **derCOP** $\equiv \mathbf{C}_{2|1}(v | u)$ and **derCOP2** $\equiv \mathbf{C}_{1|2}(u | v)$) can be written in terms of $\mathbf{C}(u | v) = \mathbf{C}_{2|1}(v | u)$ as

$$\hat{\mathbf{C}}_{2|1}(v | u) = 1 - \mathbf{C}_{2|1}(1 - v | 1 - u),$$

$$\dot{\mathbf{C}}_{2|1}(v | u) = \mathbf{C}_{2|1}(v | 1 - u), \text{ and}$$

$$\ddot{\mathbf{C}}_{2|1}(v | u) = 1 - \mathbf{C}_{2|1}(1 - v | u),$$

where the respective "surv", "acute", and "grave" are inverses (conditional quantile functions; inverses of numerical derivatives; **derCOPinv** $\equiv \mathbf{C}_{2|1}^{(-1)}(v | u)$ and **derCOPinv2** $\equiv \mathbf{C}_{1|2}^{(-1)}(u | v)$) are

$$\hat{\mathbf{C}}_{2|1}^{(-1)}(t | u) = 1 - \mathbf{C}_{2|1}^{(-1)}(1 - t | 1 - u) \rightarrow \text{"sur"},$$

$$\dot{\mathbf{C}}_{2|1}^{(-1)}(t | u) = \mathbf{C}_{2|1}^{(-1)}(t | 1 - u) \rightarrow \text{"acute"}, \text{ and}$$

$$\ddot{\mathbf{C}}_{2|1}^{(-1)}(t | u) = 1 - \mathbf{C}_{2|1}^{(-1)}(1 - t | u) \rightarrow \text{"grave"},$$

where t is a uniformly distributed variable.

To clarify the seemingly clunky nomenclature—Joe (2014) does not provide “names” for $\dot{\mathbf{C}}(u, v)$ or $\ddot{\mathbf{C}}(u, v)$ —the following guidance is informative:

- (1) "surv" or $\hat{\mathbf{C}}(u, v)$ is a reflection of U and V on the horizontal *and* vertical axes, respectively
- (2) "acute" or $\dot{\mathbf{C}}(u, v)$ is a reflection of U on the horizontal axis, and
- (3) "grave" or $\ddot{\mathbf{C}}(u, v)$ is a reflection of V on the vertical axis.

The names "acute" and "grave" match those used in the **Rd**-format math typesetting instructions.

Usage

```
simCOPmicro(u, cop=NULL, para=NULL, seed=NULL,
            reflect=c("cop", "surv", "acute", "grave",
                     "1", "2", "3", "4"), ...)
simCOPv(u, cop=NULL, para=NULL,
        reflect=c("cop", "surv", "acute", "grave",
                  "1", "2", "3", "4"), ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction. The <code>runif()</code> function in R can be used to drive conditional simulation using the <code>simCOPmicro</code> function (see Examples);
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters, if needed, to pass to the copula;
<code>seed</code>	The integer seed to pass immediately to <code>set.seed()</code> and setting it for the function <code>simCOPv</code> version will dispatch through the triple dots down to <code>simCOPmicro</code> ;
<code>reflect</code>	The reflection of the copula (see above) and the default "cop" or "1" is the usual copula definition. The numbered values correspond, respectively, to the named values; and
<code>...</code>	Additional arguments to pass should they be needed.

Value

Simulated value(s) of nonexceedance probability v are returned based on the nonexceedance probabilities u in argument `u`.

Note

The advanced features of `simCOPmicro` permit simulation of the three permutations of *variable reflection*. The first code simply produce four different "copulas" based on the *Gumbel–Hougaard* copula ($\text{GH}(u, v; \Theta)$; [GHcop](#)), which has substantial upper tail dependency but no lower tail dependency for $\Theta = 2.512$ as quantified by the [taildepCOP](#) function call.

```
U <- runif(1500); G <- 2.512; u <- 0.1; up <- 1-u; v <- 0.2; vp <- 1-v
UV <- data.frame(U, simCOPmicro(U, cop=GHcop, para=G, reflect="cop" ))
sUsV <- data.frame(U, simCOPmicro(U, cop=GHcop, para=G, reflect="surv" ))
sUV <- data.frame(U, simCOPmicro(U, cop=GHcop, para=G, reflect="acute" ))
UsV <- data.frame(U, simCOPmicro(U, cop=GHcop, para=G, reflect="grave" ))
taildepCOP(cop=GHcop, para=G) # lambdaL = 2e-05; lambdaU = 0.68224
```

The following code example will verify that the simulations produce values of U and V that are consistent with the *empirical copula* ([EMPIRCop](#)) results as well as consistent with the variable reflections provided through the [COP](#) interface. Notice the combinations of nonexceedance and exceedance probabilities blended so that the two returned values for the four different copulas are numerically congruent.

```
c(EMPIRCop(u, v, para=UV ), COP(u, v, cop=GHcop, para=G, reflect="cop" ))
c(EMPIRCop(up, vp, para=sUsV), COP(up, vp, cop=GHcop, para=G, reflect="surv" ))
c(EMPIRCop(up, v, para=sUV ), COP(up, v, cop=GHcop, para=G, reflect="acute" ))
c(EMPIRCop(u, vp, para=UsV ), COP(u, vp, cop=GHcop, para=G, reflect="grave" ))
```

The user can verify the reflections graphically using code such as this

```
xlab <- "PROBABILITY IN U"; ylab <- "PROBABILITY IN V"
layout(matrix(c(1,2,3,4), 2, 2, byrow = TRUE));
```

```

plot(UV, xlab=xlab, ylab=ylab, pch=3, lwd=0.5, col=1)
  mtext("no reflection")
plot(sUV, xlab=xlab, ylab=ylab, pch=3, lwd=0.5, col=3)
  mtext("horizontal reflection")
plot(UsV, xlab=xlab, ylab=ylab, pch=3, lwd=0.5, col=4)
  mtext("vertical reflection")
plot(sUsV, xlab=xlab, ylab=ylab, pch=3, lwd=0.5, col=2)
  mtext("double reflection")

```

in which inspection of tails exhibiting the dependency is readily seen on the four plots: upper right tail dependency (no reflection), upper left tail dependency (horizontal reflection), lower right tail dependency (vertical reflection), and lower left tail dependency (double reflection). It is important to stress that these descriptions and graphical depictions of single tail dependency are specific to the $\text{GH}(u, v; \Theta)$ copula chosen for the demonstration.

Author(s)

W.H. Asquith

References

- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
 Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[simCOP](#)

Examples

```

simCOPmicro(runif(1), cop=W ) # Frechet lower-bound copula
simCOPmicro(runif(1), cop=P ) # Independence copula
simCOPmicro(runif(1), cop=M ) # Frechet upper-bound copula
simCOPmicro(runif(1), cop=PSP) # The PSP copula

## Not run:
# Now let us get more complicated and mix two Plackett copulas together using the
# composite2COP as a "compositor." The parameter argument becomes more complex, but is
# passed as shown into composite2COP.
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop, alpha=0.3, beta=0.5, para1=0.1, para2=50)
simCOPmicro(runif(5), cop=composite2COP, para=para) #
## End(Not run)

## Not run:
# Now let us implement "our" own version of features of simCOP() but using
# the micro version to manually create just the simulation vector of V.
U <- runif(1500)
UV <- data.frame(U, simCOPmicro(U, cop=N4212cop, para=4))
plot(UV, xlab="PROBABILITY IN U", ylab="PROBABILITY IN V", pch=3, col=2) #
## End(Not run)

```

Description

Kiriliouk *et al.* (2016, pp. 360–364) describe estimation of the *spectral measure* of bivariate data. Standardize the bivariate data as X^* and Y^* as in `psepolar` and select a “large” value for the *pseudo-polar radius* S_f for nonexceedance probability f . Estimate the spectral measure $H(w)$, which is the limiting distribution of the *pseudo-polar angle* component W given that the corresponding radial component S is large:

$$\Pr[W \in \cdot | S > S_f] \rightarrow H(w) \text{ as } S_f \rightarrow \infty.$$

So, $H(w)$ is the *cumulative distribution function* of the spectral measure for angle $w \in (0, 1)$. The S_f can be specified by a nonexceedance probability F for $S_f(F)$.

The estimation proceeds as follows:

Step 1: Convert the bivariate data (X_i, Y_i) into $(\widehat{S}_i, \widehat{W}_i)$ by `psepolar` and set the threshold S_f according to “ n/k ” (this part involving k does not make sense in Kiriliouk *et al.* (2016)) where for present implementation in `copBasic` the S_f given the f by the user is based on the empirical distribution of the $\widehat{S}_i$. The empirical distribution is estimated by the *Bernstein empirical distribution* function from the `lmomco` package.

Step 2: Let I_n denote the set of indices that correspond to the observations when $\widehat{S}_i \geq S_f$ and compute N_n as the cardinality of $N_n = |I_n|$, which simply means the length of the vector I_n .

Step 3: Use the *maximum Euclidean likelihood estimator*, which is the third of three methods mentioned by Kiriliouk *et al.* (2016):

$$\widehat{H}_3(w) = \sum_{i \in I_n} \widehat{p}_{3,i} \times \mathbf{1}[\widehat{W}_i \leq w],$$

where $\mathbf{1}[\cdot]$ is an *indicator function* that is only triggered if `smooth=FALSE`, and following the notation of Kiriliouk *et al.* (2016), the “3” represents maximum *Euclidean likelihood* estimation. The $\widehat{p}_{3,i}$ are the weights

$$\widehat{p}_{3,i} = \frac{1}{N_n} [1 - (\overline{W} - 1/2) S_W^{-2} (\widehat{W}_i - \overline{W})],$$

where $\overline{W}$ is the *sample mean* and S_W^2 is the *sample variance* of $\widehat{W}_i$

$$\overline{W} = \frac{1}{N_n} \sum_{i \in I_n} \widehat{W}_i \quad \text{and} \quad S_W^2 = \frac{1}{N_n - 1} \sum_{i \in I_n} (\widehat{W}_i - \overline{W})^2,$$

where Kiriliouk *et al.* (2016, p. 363) do not show the $N_n - 1$ in the denominator for the variance but `copBasic` uses it because those authors state that the *sample variance* is used.

Step 4: A smoothed version of $\widehat{H}_3(w)$ is optionally available by

$$\tilde{H}_3(w) = \sum_{i \in I_n} \widehat{p}_{3,i} \times \mathcal{B}(w; \widehat{W}_i \nu, (1 - \widehat{W}_i) \nu),$$

where $\mathcal{B}(x; p, q)$ is the cumulative distribution function of the *Beta distribution* for $p, q > 0$ and where $\nu > 0$ is a smoothing parameter that can be optimized by cross validation.

Step 5: The *spectral density* lastly can be computed optionally as

$$\tilde{h}_3(w) = \sum_{i \in I_n} \hat{p}_{3,i} \times \beta(w; \widehat{W}_i \nu, (1 - \widehat{W}_i) \nu)$$

where $\beta(x; p, q)$ is the probability density function (pdf) of the Beta distribution. Readers are alerted to the absence of the $\mathbf{1}[\cdot]$ indicator function in the definitions of $\tilde{H}_3(w)$ and $\tilde{h}_3(w)$. This is correct and matches Kiriliouk *et al.* (2016, eqs. 17.21 and 17.22) though this author was confused for a day or so by the indicator function in what is purported to be the core definition of $\hat{H}_l(w)$ where $l = 3$ in Kiriliouk *et al.* (2016, eq. 17.21 and 17.17).

Usage

```
spectralmeas(u, v=NULL, w=NULL, f=0.90, snv=FALSE,
             smooth=FALSE, nu=100, pdf=FALSE, ...)
```

Arguments

u	Nonexceedance probability u in the X direction (actually the ranks are used so this can be a real-value argument as well);
v	Nonexceedance probability v in the Y direction (actually the ranks are used so this can be a real-value argument as well) and if NULL then u is treated as a two column R data.frame;
w	A vector of polar angle values $W \in [0, 1]$ on which to compute the $H(w)$;
f	The nonexceedance probability $F(S_f)$ of the <i>pseudo-polar radius</i> in psepolar ;
snv	Return the standard normal variate of the H by the well-known transform through the quantile function of the standard normal, <code>qnorm()</code> ;
smooth	A logical to return $\tilde{H}_3(w)$ instead of $H_3(w)$;
nu	The $\nu > 0$ smoothing parameter;
pdf	A logical to return the smoothed probability density $\tilde{h}_3(w)$. If pdf=TRUE, then internally smooth=TRUE will be set; and
...	Additional arguments to pass to the psepolar function.

Value

An R vector of $H_3(w)$, $\tilde{H}_3(w)$, or $\tilde{h}_3(w)$ is returned.

Note

The purpose of this section is to describe a CPU-intensive study of goodness-of-fit between a *Gumbel–Hougaard copula* ([GHcop](#), $\mathbf{GH}(u, v; \Theta_1)$) parent and a fitted *Hüsler–Reiss copula* ([HRcop](#), $\mathbf{HR}(u, v; \Theta_2)$). Both of these copulas are extreme values and are somewhat similar to each other, so sample sizes necessary for detection of differences should be large. A two-sided *Kolmogorov–Smirnov* tests (KS test, `ks.test()`) is used to measure significance in the differences between the estimated spectral measure distributions at $f = 0.90$ (the 90th percentile, $F(S_f)$) into the right tail.

The true copula will be the $\mathbf{GH}(\Theta_1)$ having parameter $\Theta_1 = 3.3$. The number of simulations per sample size $n \in \text{seq}(50, 1000, \text{by}=25)$ is $\text{nsim} = 500$. For each sample size, a sample from the true parent is drawn, and a $\mathbf{HR}(\Theta_2)$ fit by *maximum likelihood* (`mleCOP`). The two spectral measure distributions ($\hat{H}_{\mathbf{GH}}(w)$, H_{tru} and $\hat{H}_{\mathbf{HR}}(w)$, H_{fit}) are estimated for a uniform variate of the angle W having length equal to the applicable sample size. The KS test is made between H_{tru} and H_{fit} , and number of p-values less than the $\beta = 0.05$ (Type II error because alternative hypothesis is rigged as true) and simulation count are returned and written in file `Results.txt`. The sample sizes initially are small and traps of NaN (abandonment of a simulation run) are made. These traps are needed when the empirical distribution of H_{tru} or H_{fit} degenerates.

```
Results <- NULL
true.par <- 3.3; true.cop <- GHcop; fit.cop <- HRcop; search <- c(0,100)
nsim <- 20000; first_time <- TRUE; f <- 0.90; beta <- 0.05
ns <- c(seq(100, 1000, by=50), 1250, 1500, 1750, 2000)
for(n in ns) {
  W <- sort(runif(n)); PV <- vector(mode="numeric")
  for(i in 1:(nsim/(n/2))) {
    UV <- simCOP(n=n, cop=true.cop, para=true.par, graphics=FALSE)
    fit.par <- mleCOP(UV, cop=fit.cop, interval=search)$para
    UVfit <- simCOP(n=n, cop=fit.cop, para=fit.par, graphics=FALSE)
    Htru <- spectralmeas(UV, w=W, bound.type="Carv", f=f)
    Hfit <- spectralmeas(UVfit, w=W, bound.type="Carv", f=f)
    if(length(Htru[! is.nan(Htru)]) != length(Hfit[! is.nan(Hfit)]) |
       length(Htru[! is.nan(Htru)]) == 0 |
       length(Hfit[! is.nan(Hfit)]) == 0) {
      PV[i] <- NA; next
    } # suppressWarnings() to silence ties warnings from ks.test()
    KS <- suppressWarnings( stats::ks.test(Htru, Hfit)$p.value )
    #plot(FF, H, type="l"); lines(FF, Hfit, col=2); mtext(KS)
    message("-", i, appendLF=FALSE)
    if(length(grep("[02468]0$", as.character(i))) == 1) message()
    PV[i] <- KS
  }
  message(":", n)
  zz <- data.frame(SampleSize=n, NumPVle0p05=sum(PV[! is.na(PV)] <= beta),
                  SimulationCount=length(PV[! is.na(PV)]))
  if(first_time) { Results <- zz; first_time <- FALSE; next }
  Results <- rbind(Results, zz)
}

plot(Results$SampleSize, 100 * Results$NumPVle0p05 / Results$SimulationCount,
     type="b", cex=1.1, xlab="Sample size",
     ylab="Percent simulations with p-value < 0.05")
```

The Results show a general increase in the counts of p-value ≤ 0.05 as sample size increases. There is variation of course and increasing `nsim` would smooth that considerably. The results show for $n \approx 1,000$ that the detection of statistically significant differences for extremal $F(S_f) = 0.90$ dependency between the $\mathbf{GH}(\Theta_1=3.3)$ and $\mathbf{HR}(\Theta_2)$ are detected at the error rate implied by the specified $\beta = 0.05$.

This range in sample size can be compared to the Kullback–Leibler sample size (n_{fg}):

```
UV      <- simCOP(n=10000, cop=true.cop, para=true.par, graphics=FALSE)
fit.par <- mleCOP(UV,      cop= fit.cop, interval=search)$para
kullCOP(cop1=true.cop, para1=true.par,
        cop2=fit.cop,  para2=fit.par)$KL.sample.size
# The Kullback-Leibler (integer) sample size for detection of differences at
# alpha=0.05 are n_fg = (742, 809, 815, 826, 915, 973, 1203) for seven runs
# Do more to see variation.
```

where the *Kullback–Leibler Divergence* approach is to measure density departures across the whole $\mathcal{T}^2$ domain as opposed to extremal dependency in the right tail as does the spectral measure.

Different runs of the above code will result in different n_{fg} in part because of simulation differences internal to `kullCOP` but also because the Θ_2 has its own slight variation in its fit to the large sample simulation ($n = 10,000$) of the parent. However, it seems that $n_{fg} \approx 900$ will be on the order of the n for which the KS test on the spectral measure determines statistical significance with similar error rate.

Now, if the aforementioned simulation run is repeated for $F(S_f) = 0.95$ or $f=0.95$, the n_{fg} obviously remains unchanged at about 900 but the n for which the error rate is about $\beta = 0.05$ is $n \approx 600$. This sample size is clearly smaller than before and smaller than n_{fg} , therefore, the analysis of the empirical spectral measure deeper into the tail $F(S_f) = 0.95$ requires a smaller sample size to distinguish between the two copula—though the analysis does not address the question as to whether one or both copula are adequate for the problem at hand. For a final comparison, if the aforementioned simulation run is repeated for $F(S_f) = 0.80$ or $f=0.80$, then the n for which the error rate is about $\beta = 0.05$ is $n \approx 1,700$. Thus as analysis is made further away from the tail into the center of the distribution, the sample size to distinguish between these two similar copula increases substantially.

Author(s)

William Asquith <william.asquith@ttu.edu>

References

Kiriliouk, Anna, Segers, Johan, Warchoř, Michał, 2016, Nonparametric estimation of extremal dependence: *in* Extreme Value Modeling and Risk Analysis, D.K. Dey and Jun Yan *eds.*, Boca Raton, FL, CRC Press, ISBN 978–1–4987–0129–7.

See Also

[psepolar](#), [stabtaildepf](#)

Examples

```
## Not run:
UV <- simCOP(n=500, cop=HRCop, para=1.3, graphics=FALSE)
W <- seq(0, 1, by=0.005)
Hu <- spectralmeas(UV, w=W)
Hs <- spectralmeas(UV, w=W, smooth=TRUE, nu=100)
```

```

plot( W, Hu, type="l", ylab="Spectral Measure H", xlab="Angle")
lines(W, Hs, col=2) #
## End(Not run)

## Not run:
"GAUScop" <- function(u,v, para=NULL, ...) {
  if(length(u) == 1) u <- rep(u, length(v))
  if(length(v) == 1) v <- rep(v, length(u))
  return(copula::pCopula(matrix(c(u,v), ncol=2), para))
}
GAUSparfn <- function(rho) return(copula::normalCopula(rho, dim=2))
n <- 2000 # The PSP parent has no upper tail dependency
uv <- simCOP(n=n, cop=PSP, para=NULL, graphics=FALSE)
PLpar <- mleCOP(uv, cop=PLcop, interval=c(0,100))$para
PLuv <- simCOP(n=n, cop=PLcop, para=PLpar, graphics=FALSE)
GApafn <- mleCOP(uv, cop=GAUScop, parafn=GAUSparfn, interval=c(-1,1))$para
GAuv <- simCOP(n=n, cop=GAUScop, para=GApafn, graphics=FALSE)
GLpar <- mleCOP(uv, cop=GLcop, interval=c(0,100))$para
GLuv <- simCOP(n=n, cop=GLcop, para=GLpar, graphics=FALSE)
FF <- c(0.001, seq(0.005,0.995, by=0.005), 0.999); qFF <- qnorm(FF)
f <- 0.90 # Seeking beyond the 90th percentile pseudo-polar radius
PSPh <- spectralmeas( uv, w=FF, f=f, smooth=TRUE, snv=TRUE)
PLh <- spectralmeas(PLuv, w=FF, f=f, smooth=TRUE, snv=TRUE)
GAh <- spectralmeas(GAuv, w=FF, f=f, smooth=TRUE, snv=TRUE)
GLh <- spectralmeas(GLuv, w=FF, f=f, smooth=TRUE, snv=TRUE)
plot(qFF, PSPh, type="l", lwd=2, xlim=c(-3, 3), ylim=c(-3, 3),
     xlab="STANDARD NORMAL VARIATE OF PSEUDO-POLAR ANGLE",
     ylab="STANDARD NORMAL VARIATE OF SPECTRAL MEASURE PROBABILITY")
lines(qFF, PLh, col="red" ) # Plackett copula
lines(qFF, GAh, col="seagreen3") # Gaussian copula
lines(qFF, GLh, col="blue" ) # Galambos copula
# Notice the flat spot and less steep nature of the PSP (black line), which is
# indicative of no to even spreading tail dependency. The Plackett and Gaussian
# copulas show no specific steepening near the middle, which remains indicative
# of no tail dependency with the Plackett being less steep because it has a more
# dispersed copula density at the right tail is approached than the Gaussian.
# The Galambos copula has upper tail dependency, which is seen by
# the mass concentration and steepening of the curve on the plot.
## End(Not run)

```

Description

Kiriliouk *et al.* (2016, pp. 364–366) describe a technique for estimation of a *empirical stable tail dependence function* for a random sample. The function is defined as

$$\widehat{l}(x, y) = \frac{1}{k} \sum_{i=1}^n \mathbf{1}[R_{i,x,n} > n+1-kx \text{ or } R_{i,y,n} > n+1-ky],$$

where $\mathbf{1}[\cdot]$ is an *indicator function*, R denotes the `rank()` of the elements and $k \in [1, \dots, n]$ and k is intended to be “large enough” that $\hat{l}(x, y)$ has converged to a limit.

The “Capéraà–Fougères smooth” of the empirical stable tail dependence function is defined for a coordinate pair (x, y) as

$$\hat{l}_{CF}(x, y) = 2 \sum_{i \in I_n} \hat{p}_{3,i} \times \max[\widehat{W}_i x, (1 - \widehat{W}_i)y],$$

where $\hat{p}_{3,i}$ are the weights for the *maximum Euclidean likelihood* estimator (see [spectralmeas](#)) and $\widehat{W}_i$ are the *pseudo-polar angles* (see [spectralmeas](#)) for the index set I_n defined by $I_n = \{i = 1, \dots, n : \widehat{S}_i > \widehat{S}_{(k+1)}\}$, where $\widehat{S}_{(k+1)}$ denotes the $(k+1)$ -th largest observation of the pseudo-polar radii $\widehat{S}_i$ where the cardinality of I_n is exactly k elements long. (Tentatively, then this definition of I_n is ever so slightly different than in [spectralmeas](#).) Lastly, see the multiplier of 2 on the smooth form, and this multiplier is missing in Kiriliouk *et al.* (2016, p. 365) but shown in Kiriliouk *et al.* (2016, eq. 17.14, p. 360). Numerical experiments indicate that the 2 is needed for $\hat{l}_{CF}(x, y)$ but evidently not in $\hat{l}(x, y)$.

The visualization of $l(x, y)$ commences by setting a constant ($c > 0$) as $c_i \in 0.2, 0.4, 0.6, 0.8$ (say). The y are solved for $x \in [0, \dots, c_i]$ through the $l(x, y)$ for each of the c_i . Each solution set constitutes a *level set* for the stable tail dependence function. If the bivariate data have *asymptotic independence* (to the right), then a level set or the level sets for all the c are equal to the lines $x + y = c$. Conversely, if the bivariate data have *asymptotic dependence* (to the right), then the level sets will make 90-degree bends for $\max(x, y) = c$.

Usage

```
stabtaildepf(uv=NULL, xy=NULL, k=function(n) as.integer(0.5*n), levelset=TRUE,
             ploton=TRUE, title=TRUE, delu=0.01, smooth=FALSE, ...)
```

Arguments

uv	An R data.frame of u and v nonexceedance probabilities in the respective X (horizontal) and Y (vertical) directions. Note, <code>rank()</code> s are called on these so strictly speaking this need not be as nonexceedance probabilities. This is not an optional argument;
xy	A vector of the scalar coordinates (x, y) , which are “the relative distances to the upper endpoints of [these respective] variables” (Kiriliouk <i>et al.</i> , 2016, p. 356). This is a major point of nomenclature confusion. If these are in probability units, they are <i>exceedance probabilities</i> . Though tested for NULL and a warning issued, these can be NULL only if <code>levelset=TRUE</code> but can be set to <code>xy=NA</code> if <code>levelset=FALSE</code> and <code>smooth=TRUE</code> (see discussion in Note);
k	The k for both the $\hat{l}(x, y)$ and $\hat{l}_{CF}$, though the effect of k might not quite be the same for each. The default seems to work fairly well;
levelset	A logical triggering the construction of the level sets for $c = \text{seq}(0.1, 1, \text{by}=0.1)$;
ploton	A logical to call the <code>plot()</code> function;
title	A logical to trigger a title for the plot if <code>ploton=TRUE</code> ;
delu	The Δx for a sequence of $x = \text{seq}(0, c, \text{by}=delu)$;

smooth	A logical controlling whether $\hat{l}(x, y)$ or the Capéraà–Fougères smooth function $\hat{l}_{CF}(x, y)$ is used; and
...	Additional arguments to pass.

Value

Varies according to argument settings. In particular, the `levelset=TRUE` will cause an R list to be returned with the elements having the character string of the respective c values and the each holding a data.frame of the (x, y) coordinates.

Note

This function is also called in a secondary recursion mode. The default `levelset=TRUE` makes a secondary call with `levelset=FALSE` to compute the $\hat{l}(x, y)$ for the benefit of the looping on the one-dimensional root to solve for a single y in $\hat{l}(x, y) = c$ given a single x . If `levelset=TRUE` and `smooth=TRUE`, then a secondary call with `smooth=TRUE` and `levelset=FALSE` is made to internally return an R list containing scalar N_n and vectors $\widehat{W}_n$ and $\widehat{p}_{3,i}$ for similar looping and one-dimensional rooting for $\hat{l}_{CF}(x, y)$.

If `levelset=FALSE`, then `xy` is required to hold the (x, y) coordinate pair of interest. A demonstration follows and shows the limiting behavior of a random sample from the [N4212cop](#) copula.

```
n <- 2000 # very CPU intensive this and the next code snippet
UV <- simCOP(n=n, cop=N4212cop, para=pi); k <- 1:n
lhat <- sapply(k, function(j)
  stabtaildepf(xy=c(0.1, 0.1), uv=UV, levelset=FALSE, k=j))
plot(k, lhat, xlab="k in [1,n]", cex=0.8, lwd=0.8, type="b",
     ylab="Empirical Stable Tail Dependence Function")
mtext("Empirical function in the 0.10 x 0.10 Pr square (upper left corner)")
```

The R list that can be used to compute $\hat{l}_{CF}(x, y)$ is retrievable by

```
x <- 0.1; y <- 0.1; k <- 1:(n-1)
lhatCF <- sapply(k, function(j) {
  Hlis <- stabtaildepf(xy=NA, uv=UV, levelset=FALSE, smooth=TRUE, k=j)
  2*sum(Hlis$p3 * sapply(1:Hlis$Nn, function(i) {
    max(c(Hlis$Wn[i]*x, (1-Hlis$Wn[i])*y)) })))
})
lines(k, lhatCF, col="red")
```

The smooth line (red) of `lhatCF` is somewhat closer to the limiting behavior of `lhat`, but it is problematic to determine computational consistency. Mathematical consistency with Kiriliouk *et al.* (2016) appears to be achieved. The **Examples** section TODO.

Author(s)

William Asquith <william.asquith@ttu.edu>

References

Beirlant, J., Escobar-Bach, M., Goegebeur, Y., Guillou, A., 2016, Bias-corrected estimation of stable tail dependence function: *Journal Multivariate Analysis*, v. 143, pp. 453–466, doi:10.1016/j.jmva.2015.10.006.

Kiriliouk, Anna, Segers, Johan, Warchoř, Michał, 2016, Nonparameteric estimation of extremal dependence: *in* *Extreme Value Modeling and Risk Analysis*, D.K. Dey and Jun Yan *eds.*, Boca Raton, FL, CRC Press, ISBN 978–1–4987–0129–7.

See Also

[psepolar](#), [spectralmeas](#)

Examples

```
## Not run:
UV <- simCOP(n=1200, cop=GLcop, para=2.1) # Galambos copula
tmp1 <- stabtaildepf(UV) # the lines are curves (strong tail dependence)
tmp2 <- stabtaildepf(UV, smooth=TRUE, ploton=FALSE, col="red") #
## End(Not run)
```

statTn

The Tn Statistic of a Fitted Copula to an Empirical Copula

Description

Compute the $T_n(p)$ statistic of Genest *et al.* (2011) that is defined as

$$T_n(p) = \sum_{i=1}^n |C_n(u_i, v_i) - C_{\Theta_n}(u_i, v_i)|^p,$$

where $C_n(u, v)$ is the *empirical copula*, $C_{\Theta_n}(u, v)$ is the *fitted copula* with estimated parameters Θ_n from the sample of size n . The T_n for $p = 2$ is reported by those authors to be of general purpose and overall performance in large scale simulation studies. The extension here for arbitrary exponent p is made for flexibility. Alternatively the definition could be associated with the statistic $T_n(p)^{1/p}$ in terms of a root $1/p$ of the summation as shown above.

The T_n statistic is obviously a form of deviation between the empirical (nonparametric) and parametric fitted copula. The distribution of this statistic through Monte Carlo simulation could be used for inference. The inference is based on that a chosen parametric model is suitably close to the empirical copula. The $T_n(p)$ statistic has an advantage of being relatively straightforward to understand and explain to stakeholders and decision makers, is attractive for being suitable in a wide variety of circumstances, but intuitively might have limited statistical power in some situations for it looks at whole copula structure and not say at tail dependency. Finally, other goodness-of-fits using the squared differences between $C_n(u, v)$ and $C_{\Theta_n}(u, v)$ are [aicCOP](#), [bicCOP](#), and [rmseCOP](#).

Usage

```
statTn(u, v=NULL, cop=NULL, para=NULL, p=2, proot=FALSE, ...)
```

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction. If not given, then a second column from argument u is attempted;
cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
p	The value for p , and the default follows that of Genest <i>et al.</i> (2011);
proot	A logical controlling whether the T_n returned be rooted by $1/p$, and the default follows that of Genest <i>et al.</i> (2011); and
...	Additional arguments to pass to the copula function and (or) the empirical copula.

Value

The value for T_n is returned dependent on the specification of p and whether rooting of the result is desired.

Note

The **Examples** section shows a simple computation of the $\hat{T}_n$ statistic for a sample and a fitted copula to that sample. Ideally statTn would be wrapped in a Monte Carlo process of fitting the apparent “parent” distribution from the sample data, then for some large replication count, generate N samples of size n from the parent and from these samples compute the empirical copula and also fit parameter(s) of the chosen copula and repeatedly solve for T_n . Given a total of N values of T_n , then the sample T_n or $\hat{T}_n$ can be compared to the distribution, and if $\hat{T}_n$ is greater than say the 95th percentile, then the assumed form of the copula could be rejected.

The distTn defined below and is dependent on the `copBasic.fitpara.beta` function can be used to demonstrate concepts. (The process is complex enough that user-level implementation of distTn in **copBasic** is not presently (2019) thought appropriate.)

```
"distTn" <- function(n, N=1000, statf=NULL,
                    cop=NULL, para=para, interval=NULL, ...) {
  opts <- options(warn=-1)
  message("Estimating Tn distribution: ", appendLF=FALSE)
  Tn <- vector(mode="numeric", N)
  for(i in 1:N) {
    showi <- as.logical(length(grep("0+$", i, perl=TRUE)))
    if(showi) message(i, "-", appendLF=FALSE)
    ruv <- simCOP(n=n, cop=cop, para=para, graphics=FALSE, ...)
    rpara <- copBasic.fitpara.beta(ruv, statf=statf,
                                  interval=interval, cop=cop)
    Tn[i] <- ifelse(is.na(rpara), NA, statTn(ruv, cop=cop, para=rpara))
  }
  numNA <- length(Tn[is.na(Tn)])
  message("done: Number of failed parameter estimates=", numNA)
  options(opts)
```

```
    return(Tn[! is.na(Tn)])
  }
}
```

Let us imagine an $n = 400$ sample size of a *Galambos copula* ($\mathbf{GL}(u, v)$; `GLcop`) and then treat the *Plackett copula* ($\mathbf{PL}(u, v)$; `PLACKETTcop`) as the proper (chosen) model. The estimated parameter by the sample *Blomqvist Beta* of $\hat{\beta}_C = 0.64$ using the `blomCOP` function called from within `copBasic.fitpara.beta` is then placed in variable `para`. The $\hat{\beta}_C$ is not the most efficient estimator but for purposes here, but it is fast. The parameter for the given seed is estimated as about $\mathbf{PL}(\hat{\Theta}=20.75)$.

```
n <- 400 # sample size
correctCopula <- GLcop; set.seed(1596)
sampleUV <- simCOP(n=n, cop=correctCopula, para=1.9) # a random sample
para.correctCopula <- copBasic.fitpara.beta(uv=sampleUV, statf=blomCOP,
                                           interval=c(1,5),      cop=correctCopula)

chosenCopula <- PLACKETTcop
para <- copBasic.fitpara.beta(uv=sampleUV, statf=blomCOP,
                             interval=c(.001,200), cop=chosenCopula )
```

Next, compute the sample $\hat{T}_n = 0.063$ from `sampleUV`. The distribution of the T_n is estimated using the `distTn` function, and an estimate of the $\hat{T}_n$ p-value is in turn estimated. A large simulation run $N = 1,000$ for a sample of size of $n = 400$ is selected. The `distTn` function internally will simulate for N -replicates from the assumed parent and estimate the parameter. A computation run yields a p-value of approximately 0.01 (depending upon the seed) and is statistically significant at an alpha of 0.05, and therefore, the $\mathbf{PL}(\hat{\Theta}=20.75)$ should be rejected for fitting to these data.

```
sampleTn <- statTn(sampleUV, cop=chosenCopula, para=para)
Tns <- distTn(n=n,      cop=chosenCopula, para=para,
              interval=c(0.001, 100), statf=blomCOP)
Tns_pvalue <- 1 - sum(Tns <= sampleTn) / length(Tns) # estimate p-value
```

The demonstration is furthered with a check on the *Kullback–Leibler sample size* n_{fg} at the 5-percent significance level ($\alpha = 0.05$) by the `kullCOP` function, which yields 100. Given the parent copula as $\mathbf{GL}(\Theta=1.9)$, therefore, it would take approximately 100 samples to distinguish between that copula and a $\mathbf{PL}(\Theta=20.75)$ where in this case the fit was through the $\hat{\beta}_C = 0.64$.

```
kullCOP(cop1=correctCopula, para1=1.9,
        cop2=chosenCopula, para2=para)$KL.sample.size # KL sample size = 100
vuongCOP(sampleUV, cop1=correctCopula, para1=para.correctCopula,
          cop2=chosenCopula, para2=para)$message
# [1] "Copula 1 has better fit than Copula 2 at 100x(1-alpha) level"
```

The available sample size $n = 400$ is then about four times larger than n_{fg} so the sample size n should be sufficient to judge goodness-of-fit. This is a large value but with the sample variability of $\hat{\beta}_C$, it seems that other measures of association such as *Spearman Rho* (`rhoCOP`) or *Kendall Tau* (`tauCOP`) and others cross-referenced therein might be preferable.

The prior conclusion is supported by the p-value of the $\hat{T}_n$ being about 0.01, which suggests that the $\mathbf{PL}(u, v)$ is not a good model of the available sample data in `sampleUV`. Lastly, these judgments

are consistent with the *Vuong Procedure* performed by the `vuongCOP` function, which reports at the 5-percent significance level that “copula number 1”—in this case, the $\mathbf{GL}(u, v)$ —has the better fit, and this is obviously consistent with the problem setup because the random sample for investigation was drawn from the Galambos coupla (the parent form).

Author(s)

W.H. Asquith

References

Genest, C., Kojadinovic, I., Nešlehová, J., and Yan, J., 2011, A goodness-of-fit test for bivariate extreme-value copulas: Bernoulli, v. 17, no. 1, pp. 253–275.

See Also

`aicCOP`, `bicCOP`, `rmseCOP`, `vuongCOP`, `kullCOP`

Examples

```
## Not run:
# Example here is just for Tn. For the example below, the PSP copula is quite different
# from the Gumbel-Hougaard copula and thus, the hatTn would be expected to be different
# from those of the Gumbel-Hougaard and certainly not too near to zero.
samUV <- simCOP(n=60, cop=PSP, graphics=FALSE, seed=1) # random sample
hatTau <- cor(samUV$U, samUV$V, method="kendall") # Kendall Tau
hatTn <- statTn(samUV, cop=GHCop, para=GHCop(tau=hatTau)$para,
               ctype="bernstein", bernprogress=TRUE) # 0.03328789
# hatTn in this case is by itself is somewhat uninformative and requires
# Monte Carlo to put an individual value into context.
## End(Not run)
```

surCOP

The Survival Copula

Description

Compute the *survival copula* from a copula (Nelsen, 2006, pp. 32–34), which is defined as

$$\hat{\mathbf{C}}(1 - u, 1 - v) = \hat{\mathbf{C}}(u', v') = \Pr[U > u, V > v] = u' + v' - 1 + \mathbf{C}(1 - u', 1 - v'),$$

where u' and v' are exceedance probabilities and $\mathbf{C}(u, v)$ is the copula (`COP`). The *survival copula* is a reflection of both U and V .

The *survival copula* is an expression of the joint probability that both $U > v$ and $U > v$ when the arguments a and b to $\hat{\mathbf{C}}(a, b)$ are exceedance probabilities as shown. This is unlike a copula that has $U \leq u$ and $V \leq v$ for nonexceedance probabilities u and v . Alternatively, the joint probability that both $U > u$ and $V > v$ can be solved using just the copula $1 - u - v + \mathbf{C}(u, v)$, as shown

below where the arguments to $\mathbf{C}(u, v)$ are nonexceedance probabilities. The later formula is the *joint survival function* $\overline{\mathbf{C}}(u, v)$ ([surfuncCOP](#)) defined for a copula (Nelsen, 2006, p. 33) as

$$\overline{\mathbf{C}}(u, v) = \Pr[U > u, V > v] = 1 - u - v + \mathbf{C}(u, v).$$

Users are directed to the collective documentation in [COP](#) and [simCOPmicro](#) for more details on copula reflection.

Usage

```
surCOP(u, v, cop=NULL, para=NULL, exceedance=TRUE, ...)
```

Arguments

u	Exceedance probability $u' = 1 - u$ (u nonexceedance based on exceedance) in the X direction;
v	Exceedance probability $v' = 1 - v$ (v nonexceedance based on exceedance) in the Y direction;
cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
exceedance	A logical affirming whether u and v are really in exceedance probability or not? If FALSE, then the complements of the two are made internally and the nonexceedances can thus be passed; and
...	Additional arguments to pass (such as parameters, if needed, for the copula in the form of an R list).

Value

Value(s) for the survival copula are returned.

Note

The author (Asquith) finds the use of exceedance probabilities delicate in regards to Nelsen's notation. This function and [coCOP](#) have the exceedance argument to serve as a reminder that the survival copula as usually defined uses exceedance probabilities as its arguments.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[COP](#), [coCOP](#), [duCOP](#), [surfuncCOP](#), [simCOPmicro](#)

Examples

```

u <- 0.26; v <- 0.55 # nonexceedance probabilities
up <- 1 - u; vp <- 1 - v # exceedance probabilities
surCOP(up, vp, cop=PSP, exceedance=TRUE) # 0.4043928
surCOP(u, v, cop=PSP, exceedance=FALSE) # 0.4043928 (same because of symmetry)
surfuncCOP(u, v, cop=PSP) # 0.4043928
# All three examples show joint prob. that U > u and V > v.

## Not run:
# A survival copula is a copula so it increases to the upper right with increasing
# exceedance probabilities. Let us show that by hacking the surCOP function into
# a copula for feeding back into the algorithmic framework of copBasic.
UsersCop <- function(u,v, para=NULL) {
  afunc <- function(u,v, theta=para) { surCOP(u, v, cop=N4212cop, para=theta)}
  return(asCOP(u,v, f=afunc)) }
image(gridCOP(cop=UsersCop, para=1.15), col=terrain.colors(20),
      xlab="U, EXCEEDANCE PROBABILITY", ylab="V, EXCEEDANCE PROBABILITY") #
## End(Not run)

```

surfuncCOP

*The Joint Survival Function***Description**

Compute the *joint survival function* for a copula (Nelsen, 2006, p. 33), which is defined as

$$\overline{\mathbf{C}}(u, v) = \Pr[U > u, V > v] = 1 - u - v + \mathbf{C}(u, v) = \hat{\mathbf{C}}(1 - u, 1 - v),$$

where $\hat{\mathbf{C}}(u', v')$ is the *survival copula* ([surCOP](#)), which is defined by

$$\hat{\mathbf{C}}(u', v') = \Pr[U > u, V > v] = u' + v' - 1 + \mathbf{C}(1 - u', 1 - v').$$

Although the joint survival function is an expression of the probability that both $U > v$ and $U > v$, $\overline{\mathbf{C}}(u, v)$ is not a copula.

Usage

```
surfuncCOP(u, v, cop=NULL, para=NULL, ...)
```

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction;
cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula; and
...	Additional arguments to pass (such as parameters, if needed, for the copula in the form of a list.

Value

Value(s) for the joint survival function are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[surCOP](#)

Examples

```
"M0cop.formula" <- function(u,v, para=para, ...) {
  alpha <- para[1]; beta <- para[2]; return(min(v*u^(1-alpha), u*v^(1-beta)))
}
"M0cop" <- function(u,v, ...) { asCOP(u,v, f=M0cop.formula, ...) }
u <- 0.2; v <- 0.75; ab <- c(1.5, 0.3)
# U **and** V are less than or equal to a threshold +
# U **or** V are less than or equal to a threshold
surCOP(1-u,1-v, cop=M0cop, para=ab) + duCOP(u,v, cop=M0cop, para=ab) # UNITY
surfuncCOP(u,v, cop=M0cop, para=ab) + duCOP(u,v, cop=M0cop, para=ab) # UNITY

## Not run:
# The joint survival function is not a copula. So, it does not increases to the upper
# right with increasing exceedance probabilities. Let us show that by hacking the surCOP
# function into a copula for feeding back into the algorithmic framework of copBasic.
UsersCop <- function(u,v, para=NULL) {
  afunc <- function(u,v, theta=para) { surfuncCOP(u, v, cop=N4212cop, para=theta) }
  return(asCOP(u,v, f=afunc)) }
image(gridCOP(cop=UsersCop, para=1.15), col=terrain.colors(20),
      xlab="U, NONEXCEEDANCE PROBABILITY", ylab="V, NONEXCEEDANCE PROBABILITY") #
## End(Not run)

## Not run:
# Conditional return period (Salvadori et al., 2007, p. 159)
UV <- simCOP(n=100000, cop=PLACKETTcop, para=5, graphics=FALSE)
u <- 0.5; v <- 0.99; cd <- UV$V[UV$U > u]
by.counting <- length(cd[cd > v]) / length(cd) # 0.0172
by.theo <- surfuncCOP(u,v, cop=PLACKETTcop, para=5) / (1-u) # 0.0166
by.ec <- surfuncCOP(u,v, cop=EMPIRCop, para=UV) / (1-u) # 0.0189
print(1/by.theo) # conditional return period for V > 0.99 given U > 0.5
## End(Not run)
```

Description

Compute the *tail concentration function* (q_C) of a copula $C(u, v)$ ([COP](#)) or diagonal ([diagCOP](#)) of a copula $\delta_C(t) = C(t, t)$ according to Durante and Semp (2015, p. 74):

$$q_C(t) = \frac{C(t, t)}{t} \cdot \mathbf{1}_{[0, 0.5)} + \frac{1 - 2t + C(t, t)}{1 - t} \cdot \mathbf{1}_{[0.5, 1]} \quad \text{or}$$

$$q_C(t) = \frac{\delta_C(t)}{t} \cdot \mathbf{1}_{[0, 0.5)} + \frac{1 - 2t + \delta_C(t)}{1 - t} \cdot \mathbf{1}_{[0.5, 1]},$$

where t is a nonexceedance probability on the margins and $\mathbf{1}(\cdot)$ is an *indicator function* scoring 1 if condition is true otherwise zero on what interval t resides: $t \in [0, 0.5)$ or $t \in [0.5, 1]$. The $q_C(t; \mathbf{M}) = 1$ for all t for the [M](#) copula and $q_C(t; \mathbf{W}) = 0$ for all t for the [W](#) copula. Lastly, the function is related to the *Blomqvist Beta* (β_C ; [blomCOP](#)) by

$$q_C(0.5) = (1 + \beta_C)/2,$$

where $\beta_C = 4C(0.5, 0.5) - 1$. Lastly, the $q_C(t)$ for $0, 1 = t$ is NaN and no provision for alternative return is made. Readers are asked to note some of the mathematical similarity in this function to Blomqvist Betas in [blomCOPss](#) in regards to tail dependency.

Usage

```
tailconCOP(t, cop=NULL, para=NULL, ...)
```

Arguments

<code>t</code>	Nonexceedance probabilities t ;
<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula; and
<code>...</code>	Additional arguments to pass to the copula function.

Value

Value(s) for q_C are returned.

Author(s)

W.H. Asquith

References

Durante, F., and Sempi, C., 2015, Principles of copula theory: Boca Raton, CRC Press, 315 p.

See Also

[taildepCOP](#), [tailordCOP](#)

Examples

```
tailconCOP(0.5, cop=PSP) == (1 + blomCOP(cop=PSP)) / 2 # TRUE
```

taildepCOP

The Lower- and Upper-Tail Dependency Parameters of a Copula

Description

Compute the *lower- and upper-tail dependency parameters* (if they exist), respectively, of a copula according to Nelsen (2006, pp. 214–215). Graphical confirmation of the computations is important, and therefore, the function can also generate a plot. The dependency parameters are expressions of conditional probability that Y is greater than the $100 \times t$ -th percentile of its distribution G given that X is greater than the $100 \times t$ -th percentile of its distribution F as t approaches unity. Specifics in terms of quantile functions $G^{(-1)}(t) = y(t)$ and $F^{(-1)}(t) = x(t)$ follow.

The *lower-tail dependence parameter* $\lambda_{\mathbf{C}}^L$ is defined as

$$\lambda_{\mathbf{C}}^L = \lim_{t \rightarrow 0^+} \Pr[Y \leq y(t) \mid X \leq x(t)], \text{ and}$$

the *upper-tail dependence parameter* $\lambda_{\mathbf{C}}^U$ with reversed inequalities is defined as

$$\lambda_{\mathbf{C}}^U = \lim_{t \rightarrow 1^-} \Pr[Y > y(t) \mid X > x(t)].$$

Nelsen (2006, p. 214) also notes that both $\lambda_{\mathbf{C}}^L$ and $\lambda_{\mathbf{C}}^U$ are nonparametric and depend only on the copula of X and Y , and Nelsen shows that each can be computed if the above limits exist as follows:

$$\lambda_{\mathbf{C}}^L = \lim_{t \rightarrow 0^+} \frac{\mathbf{C}(t, t)}{t} = \delta'_{\mathbf{C}}(0^+) \text{ and}$$

$$\lambda_{\mathbf{C}}^U = \lim_{t \rightarrow 1^-} \frac{1 - 2t - \mathbf{C}(t, t)}{1 - t} = 2 - \lim_{t \rightarrow 1^-} \frac{1 - \mathbf{C}(t, t)}{1 - t} = 2 - \delta'_{\mathbf{C}}(1^-),$$

where $\delta'_{\mathbf{C}}(t)$ is the derivative of the diagonal of the copula. Multiple presentations are shown because algebraic variants are shown across the literature.

If $\lambda_{\mathbf{C}}^L \in (0, 1]$, then $\mathbf{C}$ has lower-tail dependence but if $\lambda_{\mathbf{C}}^L = 0$, then $\mathbf{C}$ has *no* lower-tail dependence. Likewise, if $\lambda_{\mathbf{C}}^U \in (0, 1]$, then $\mathbf{C}$ has upper-tail dependence but if $\lambda_{\mathbf{C}}^U = 0$, then $\mathbf{C}$ has *no* upper-tail dependence.

Usage

```
taildepCOP(cop=NULL, para=NULL, tol=1e-6, divisor=2, plot=FALSE, ylim=NULL,
           verbose=FALSE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
tol	A tolerance on convergence;
divisor	The divisor on the incremental reductions towards 0^+ and 0^- by the algorithm;
plot	A logical plotting a diagnostic plot of the diagonal derivatives and label the limits;
ylim	Optional vertical limits if the plot is turned on. Although the dependence parameters are bounded as described above, numerical stability can be a problem. Stability is especially a problem if an empirical copula is being used; therefore, the bounds of the plot are left open unless the user locks them down with this argument;
verbose	Show incremental progress; and
...	Additional arguments to pass to the copula function.

Value

An R list is returned.

lambdaL	The rounded value of λ_G^L ;
lambdaU	The rounded value of λ_G^U ;
source	An attribute identifying the computational source: “taildepCOP”.

Note

IMPLEMENTED ALGORITHM—The algorithm implemented for taildepCOP is based on halves (or alternatives by the setting of divisor argument) and uses the copula function (not an analytical or even numeric derivative of the diagonal, $\delta'_G(t)$). Starting from the median or $t = 0.5$, each limit is respectively computed by successive halving (or the setting of argument divisor) of the distance towards 0^+ and 1^- and checking the change in computed value against the tolerance tol argument. After the change becomes less than the tolerance, convergence is assumed. Other tests are made for NaN to aid in breaking the successive halvings. The rounding for the numerical results for λ_G^U and λ_G^L is an order of magnitude larger than the tolerance.

Users are encouraged to plot the results and further verify whether the convergence makes sense. The plot produced when plot=TRUE shows the probability t transformed into standard normal variates by the qnorm() function in R so that the distal reaches of each tail and thus limit are readily seen. The terminal points of each limit computation are shown by a small dot and the letter “L” and “U” also are plotted at the terminal points.

Joe (2014, p. 63) reports that “the empirical measure of tail dependence $[\hat{\lambda}_G^L \text{ or } \hat{\lambda}_G^U]$ for data does not really exist because of the limit.” Joe (2014) suggests that other sources in the literature are the “best that can be done” (Dobrić and Schmid, 2005; Frahm *et al.*, 2005). Another source of discussion is by Schmidt and Stadtmüller (2006). The results therein are not yet followed up for the **copBasic** package. Picking up the simulation dealt with extensively in the **Note** section of [vuongCOP](#), a user might try this:

```

set.seed(385); n <- 390
UV <- simCOP(cop=PSP, n=n, col=8, pch=16, graphics=FALSE)
taildepCOP(cop=EMPIRCop, para=UV, plot=TRUE, divisor=8, ylim=c(0,1))
taildepCOP(cop=PSP) # lower=0.5, upper=0

```

The returned tail dependency parameters are numerically of little importance and in strict terms likely misleading. What should be of interest are the plotted trajectories of the lower and upper lines. Note: the lower tail wobbles but seems to show stability towards near $\hat{\lambda}_G^L = 0.5$ and upper-tail line wobbles downward towards $\hat{\lambda}_G^U = 0$. These values are, respectively, the tail dependencies of the **PSP** copula ([PSP](#)). A user might try increasing the sample size by an order of magnitude and rerunning the above code. Lastly, Salvadori *et al.* (2006, pp. 173–175) caution on the difficulties of nonparametric tail dependency estimation. Given objectives of the **copBasic** package, estimation of $\hat{\lambda}_G^L$ and $\hat{\lambda}_G^U$, therefore, is an open development opportunity.

DEMONSTRATION (Tail Dependence)—The following example shows a comparison between early code examples by Charpentier (2012) concerning copulas and tail dependence using real-world data. Consider the lossalae data set and the following code requiring the **evd** package:

```

library(evd); X <- lossalae # Charpentier (2012)
library(copBasic)
fakeU <- lmomco::pp(X[,1],sort=FALSE,a=0) # Weibull plotting position i/(n+1)
fakeV <- lmomco::pp(X[,2],sort=FALSE,a=0) # Weibull plotting position i/(n+1)
uv <- data.frame(U=fakeU, V=fakeV) # parameter "object" for Empirical copula
plot(uv)
TD <- taildepCOP(cop=EMPIRCop, para=uv, divisor=25,
                 plot=TRUE, ylim=c(0, 7/10))
U <- rank(X[,1])/(nrow(X)+1); V <- rank(X[,2])/(nrow(X)+1) # Charpentier(2012)
Lemp <- function(z) sum((U<=z) & (V<=z)) / sum(U<=z) # Charpentier(2012)
Remp <- function(z) sum((U>=1-z) & (V>=1-z)) / sum(U>=1-z) # Charpentier(2012)
u <- seq(0.001, 0.5, by=.001) # Charpentier(2012)
L <- Vectorize(Lemp)(u); R <- Vectorize(Remp)(rev(u)) # Charpentier(2012)
lines(qnorm(c(u, u + 0.5 - u[1])), c(L,R)) # modified after Charpentier(2012)
legend("bottomright", c("Lower-tail dependency by taildepCOP()",
                        "Upper-tail dependency by taildepCOP()",
                        "Charpentier (2012)"), bty="n", cex=0.9,
      lwd=1, lty=1, col=c("red", "blue", "black"))

```

The figure that will have been generated shows considerably similarity to that from the algorithms of Charpentier. Now, let us extend the discussion by using the *Blomqvist (Schmid-Schmidt) Betas* ([blomCOPss](#)) that have a formulation permitting lower- and upper-tail dependency parameters in a different manner than the definitions of this documentation for taildepCOP.

```

edge <- 30 * 1 / (1+nrow(X)) # as few as 30 samples into the tails
psl <- pnorm(seq(0, qnorm(edge), by=-0.005))
psu <- pnorm(seq(0, qnorm(1-edge), by= 0.005))
lines(qnorm(psl),
      sapply(psl, function(p) { blomCOPss(as.sample=TRUE, para=uv,
      ctype="checkerboard", uu=rep(p, 2), vv=c(1,1)) })),
      col="darkgreen", lty=1, lwd=2)

```

```

lines(qnorm(psu),
      sapply(psu, function(p) { blomCOPss(as.sample=TRUE, para=uv,
      ctype="checkerboard", uu=c(0,0), vv=rep(p, 2)) }),
      col="darkgreen", lty=1, lwd=2)
points(0, blomCOP(as.sample=TRUE, para=uv), pch=16, col="magenta", cex=2)
legend("topleft", c("Tail dependency by Blomqvist (Schmid-Schmidt) Betas",
      "Blomqvist Beta C(1/2, 1/2)",
      bty="n", cex=0.9, lwd=2, lty=c(1,NA), pch=c(NA,16),
      pt.cex=c(NA,2), col=c("darkgreen", "magenta"))

```

The thick green lines show that the dependency parameters to the left and right are approached along a different trajectory using the definitions in `blomCOPss`. It seems at some stage in analysis that the practitioner will need to decide how deep into the tail the sample will permit for a reliable estimate of the dependency parameters. *Blomqvist Beta* ($\hat{\beta}_C$) (`blomCOP` is shown as the magenta dot at the median and henceforth from do the trajectories of $\hat{\lambda}_{\beta_C}^L$ and $\hat{\lambda}_{\beta_C}^U$ extend as their $\lim_{p \rightarrow 0^+}$. Ultimately, for the data shown in the figure, perhaps the $\hat{\lambda}_C^L = 0.1$ and the $\hat{\lambda}_C^U = 0.3$ and a parametric copula fitted in part to such values.

Author(s)

W.H. Asquith

References

- Charpentier, A., 2012, Copulas and tail dependence, part 1: R-bloggers, dated Sept. 17, 2012, accessed on February 2, 2019 at <https://www.r-bloggers.com/2012/09/copulas-and-tail-dependence-part-1/>
- Dobrić, J. and Schmid, F., 2005, Nonparametric estimation of the lower tail dependence λ^L in bivariate copulas: Journal of Applied Statistics, v. 32, no. 4, pp. 387–407, doi:10.1080/02664760500079217.
- Frahm, G., Junker, M., and Schmidt, R., 2005, Estimating the tail-dependence coefficient—Properties and pitfalls: Insurance—Mathematics and Economics, v. 37, no. 1, pp. 80–100.
- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, Extremes in Nature—An approach using copulas: Springer, 289 p.
- Schmidt, R., and Stadtmüller, U., 2006, Nonparametric estimation of tail dependence: The Scandinavian Journal of Statistics, v. 33, pp. 307–335.

See Also

`COP`, `tailconCOP`, `tailordCOP`, `blomCOPss`

Examples

```

# Plot the tail dependencies by nonexceedance probability for a
# for a positive association Plackett copula and see that both are zero.

```

```

taildepCOP(cop=PLACKETTcop, para=3, plot=TRUE)
# So, Plackett has no tail dependency, as Nelsen (2006, p. 215) shows.

## Not run:
# The results that follow match those reported by Nelsen (2006, p. 215).
taildepCOP(cop=M0cop, para=c(0.4, 0.9)) # LambL = 0, LambU = 0.4 [min(alpha,beta)]
## End(Not run)

## Not run:
# Analytical solution to Gumbel-Hougaard copula from the copula package:
copula::lambda(copula::gumbelCopula(3))
# lower upper
# 0.000000 0.740079
# Numerical approximation (see copBasic::GHcop for analytical formula):
as.data.frame(taildepCOP(GHcop, para=3))
# lambdaL lambdaU source
#1 0.00012 0.74008 taildepCOP
## End(Not run)

## Not run:
# Plot the tail dependencies by nonexceedance probability
# for the PSP copula, which has lower but no upper-tail dependence.
taildepCOP(cop=PSP, para=NULL, plot=TRUE) # LambL=0.5, LambU=0
# which is readily confirmed by simCOP(1000, cop=PSP)
# Nelsen (2006, p. 216) reports that this copula has LambL=1/2 and LambU=0,
# and we get the same results here.

# How about some composited Plackett-Plackett copulas?
# Each has upper- and lower-tail dependence parameters equal to zero.
para <- list( cop1=PLACKETTcop, cop2=PLACKETTcop, alpha=0.9392,
              para1=0.00395, para2=4.67, beta=0.5699)
taildepCOP(cop=composite2COP, para=para, plot=TRUE, verbose=TRUE) #
## End(Not run)

## Not run:
# This next Plackett-Plackett is interesting because at its core it looks
# like it should be both tail dependent like M() but the shapes of the curves
# are quite different from those of M(). This example shows numerical
# instability for the upper tail but not the lower tail. So, we extend the
# example to shown the tail dependency trajectories by blomCOPss(). And again
# it is seen that the lower tail as a stable solution but the upper tail
# has instability at 6 standard deviations into the upper tail.
para <- list( cop1=PLACKETTcop, cop2=PLACKETTcop, alpha=0.0063,
              para1=0.101, para2=4493, beta=0.0167)
taildepCOP(cop=composite2COP, para=para, plot=TRUE)
lsu <- pnorm(seq(-7, 0, by=.01))
psu <- pnorm(seq( 0, 7, by=.01))
lines(qnorm(lsu), sapply(lsu, function(p) {
  blomCOPss(cop=composite2COP, para=para, vv=c(1,1), uu=rep(p, 2)) }},
      col="darkgreen", lty=2, lwd=1)
lines(qnorm(psu), sapply(psu, function(p) {
  blomCOPss(cop=composite2COP, para=para, uu=c(0,0), vv=rep(p, 2)) }},
      col="darkgreen", lty=2, lwd=1) #

```

```
## End(Not run)
```

tailordCOP

The Lower- and Upper-Tail Orders of a Copula

Description

Compute the *lower-* and *upper-tail orders* (if they exist), respectively, of a copula $\mathbf{C}(u, v)$ according to Joe (2014, pp. 67–70). The *tail order* is a concept for the strength of dependence in the joint tails of a multivariate distribution. The opposing tails can be compared to assess tail order or *reflection symmetry* (term by Joe (2014) for Nelsen’s (2006, p. 36) term *radial symmetry*). Joe (2014) provides extensively analytical details but sufficient for the **copBasic** package, the tail orders can be numerically explored.

The *lower-tail order* maybe numerically approximated by

$$\kappa_{\mathbf{C}}^L = \frac{\log(\mathbf{C}(t, t))}{\log(t)},$$

for some small positive values of t , and similarly the *upper-tail order* maybe numerically approximated by

$$\kappa_{\mathbf{C}}^U = \frac{\log(\hat{\mathbf{C}}(t, t))}{\log(t)},$$

where $\hat{\mathbf{C}}(u, v)$ is the *survival copula* ([surCOP](#)). Joe (2014) has potentially(?) conflicting notation in the context of the upper-tail order; the term “reflection” is used (p. 67) and “lower tail order of the reflected copula is the same as the upper tail order of the original copula” (p. 69), but Joe (2014, p. 67) only uses the joint survival function ([surfuncCOP](#)) in the definition of $\kappa_{\mathbf{C}}^U$.

As a note, the author of this package was not able to get tailordCOP to function properly for the upper-tail order using the joint survival function as implied on the bottom of Joe (2014, p. 67) and fortunately the fact that “reflection” is used in other contexts and used in analytical examples, the tailordCOP function uses the lower-tail order of the reflection (survival copula). Joe (2014) also defines *tail order parameter* Ψ but that seems to be a result of analytics and not implemented in this package. Lastly, the tail orders are extendable into d dimensions, but only a bivariate ($d = 2$) is provided in **copBasic**. The tail orders have various classifications for $\kappa = \kappa_L = \kappa_U$:

- *Intermediate tail dependence* for $1 < \kappa < d$ or $\kappa = 1, \Psi = 0$;
- *Strong tail dependence* for $\kappa = 1$ with $\Psi > 0$; and
- *Tail orthant independence* or *tail quadrant independence* for $\kappa = d$.

Joe (2014) provides additional properties:

- $\kappa_L = \kappa_U = d$ for the d -dimensional *independence copula* ([P](#); e.g. tailordCOP(cop=P));
- It is not possible for $\kappa_L < 1$ or $\kappa_U < 1$ but each can be > 1 for a $\mathbf{C}(u, v)$ having some negative dependence (e.g. tailordCOP(cop=PLACKETTcop, para=0.2); see [PLACKETTcop](#)); and
- For the bivariate *Fréchet–Hoeffding lower-bound copula* ([W](#); *countermonotonicity copula*) the $\kappa_L = \kappa_U$ and can be considered $+\infty$. (A special trap in the tailordCOP provides consistency on [W](#) but does not test that the copula is actually that function itself.)

Usage

```
tailordCOP(cop=NULL, para=NULL, tol=1e-6, plot=FALSE, verbose=FALSE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
tol	A tolerance on convergence;
plot	A logical plotting a diagnostic plot of the diagonal derivatives and label the limits;
verbose	Show incremental progress; and
...	Additional arguments to pass to the copula function.

Value

An R list is returned.

kappaL	The rounded value of κ_C^L ;
kappaU	The rounded value of κ_C^U ;
source	An attribute identifying the computational source: “tailordCOP”.

Note

The algorithm implemented for tailordCOP is based on halves (or alternatives by the setting of the divisor argument) and uses the copula function (not an analytical or even numerical derivative of the diagonal, $\delta'_C(t)$). Starting from the median or $t = 0.5$, each limit is respectively computed by successive halving of the distance towards 0^+ and checking the change in computed value against the tolerance tol argument. After the change becomes less than the the tolerance, convergence is assumed. Other tests are made for NaN to aid in breaking the successive halving. The rounding for the numerical results for κ_C^U and κ_C^L is an order of magnitude larger than the tolerance.

Users are encouraged to plot the results and further verify whether the convergence makes sense. The plot produced when plot=TRUE shows the probability t transformed into standard normal variates by the qnorm() function in R so that the distal reaches of each tail and thus limit are readily seen. The terminal points of each limit computation are shown by a small dot, and the letter “L” and “U” also are plotted at the terminal points.

Author(s)

W.H. Asquith

References

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also

[COP](#), [tailconCOP](#), [taildepCOP](#)

Examples

```
## Not run:
# Joe (2014, p. 5) names MTCJ = Mardia-Takahasi-Cook-Johnson copula
"MT CJ" <- function(u,v, para) { (u^(-para) + v^(-para) - 1)^(-1/para) }
# The results that follow match those reported by Joe (2014, p. 69) who
# analytically derives KappaL = 1 and KappaU = 2.
# TAIL ORDER:
tailordCOP(cop=MTCJ, para=3, plot=TRUE) # kappaL = 1.00667, kappaU = 1.96296
# TAIL DEPENDENCY:
taildepCOP(cop=MTCJ, para=3, plot=TRUE) # lambdaL = 0, lambdaU = 0.7937
# Joe (2014) reports lambdaL = 2^(-1/para) = 2^(-1/3) = 0.7937005
## End(Not run)
```

tauCOP

The Kendall Tau and Concordance Function of a Copula

Description

Compute the measure of association known as the *Kendall Tau* (τ_C) of a copula (τ_C) according to Nelsen (2006, sec. 5.1.1 and p. 161) by

$$\tau_C = \mathcal{Q}(\mathbf{C}, \mathbf{C}) = 4 \int \int_{\mathcal{T}^2} \mathbf{C}(u, v) d\mathbf{C}(u, v) - 1,$$

where $\mathcal{Q}(\mathbf{C}, \mathbf{C})$ is a *concordance function* (concordCOP) of a copula with itself. Nelsen (2006, p. 164) reports however that this form is often not amenable to computation when there is a singular component to the copula and that the expression

$$\tau_C = 1 - 4 \int \int_{\mathcal{T}^2} \frac{\delta \mathbf{C}(u, v)}{\delta u} \frac{\delta \mathbf{C}(u, v)}{\delta v} du dv$$

is to be preferred. Such an expression hence relies on the partial numerical derivatives of the copula provided by [derCOP](#) and [derCOP2](#). The Nelsen (2006) preferred expression is used by the tauCOP function. Nelsen (2006, pp. 175–176) reports that the relation between τ_C and ρ_C ([rhoCOP](#)) is $-1 \leq 3\tau - 2\rho \leq 1$ (see [rhoCOP](#) for more details).

Nelsen (2006, pp. 160–161) lists some special identities involving $\mathcal{Q}(\mathbf{C}_1, \mathbf{C}_2)$:

$$\mathcal{Q}(\mathbf{M}, \mathbf{M}) = 4 \int_0^1 u du - 1 = 1,$$

$$\mathcal{Q}(\mathbf{M}, \mathbf{\Pi}) = 4 \int_0^1 u^2 du - 1 = 1/3,$$

$$\mathcal{Q}(\mathbf{M}, \mathbf{W}) = 4 \int_{1/2}^1 (2u - 1) du - 1 = 0,$$

$$\mathcal{Q}(\mathbf{W}, \mathbf{\Pi}) = 4 \int_0^1 u(1 - u) du - 1 = -1/3,$$

$$\mathcal{Q}(\mathbf{W}, \mathbf{W}) = 4 \int_0^1 0 \, du - 1 = -1, \text{ and}$$

$$\mathcal{Q}(\mathbf{\Pi}, \mathbf{\Pi}) = 4 \iint_{T^2} uv \, du dv - 1 = 0.$$

Kendall Tau also can be expressed in terms of the *Kendall Function* ($F_K(z)$; [kfuncCOP](#)):

$$\tau_C = 3 - 4 \int_0^1 F_K(t) \, dt,$$

which is readily verified by code shown in **Examples**. This definition might be useful if integration errors are encountered for some arbitrary copula and arbitrary parameter set. In fact, should two attempts (see source code) at dual integration of the partial derivatives occur, the implementation switches over to integration of the Kendall Function (*e.g.* `tauCOP(cop=N4212cop, para=2)`). Note, Durante and Sempi have erroneously dropped the multiplication by “4” as shown above in their definition of τ_C as a function of $F_K(t)$ (Durante and Sempi, 2015, eq. 3.9.4, p. 121).

Usage

```
tauCOP( cop=NULL, para=NULL,
        cop2=NULL, para2=NULL, as.sample=FALSE, brute=FALSE, delta=0.002, ...)

concordCOP(cop=NULL, para=NULL, cop2=NULL, para2=NULL, ...)
```

Arguments

<code>cop</code>	A copula function;
<code>para</code>	Vector of parameters or other data structure, if needed, to pass to the copula;
<code>cop2</code>	A second copula function;
<code>para2</code>	Vector of parameters or other data structure, if needed, to pass to the second copula;
<code>as.sample</code>	A logical controlling whether an optional <code>R</code> <code>data.frame</code> in <code>para</code> is used to compute the $\hat{\tau}$ by dispatch to <code>cor()</code> function in <code>R</code> with <code>method = "kendall1"</code> ;
<code>brute</code>	Should brute force be used instead of two nested <code>integrate()</code> functions in <code>R</code> to perform the double integration;
<code>delta</code>	The du and dv for the brute force integration using <code>brute</code> ; and
<code>...</code>	Additional arguments to pass on to derCOP and derCOP2 .

Value

The value for τ_C is returned.

Note

Although titled for computation of the Kendall Tau, the tauCOP function also is the implementation of the *concordance function* $\mathcal{Q}(\mathbf{C}_1, \mathbf{C}_2)$ (see Nelsen (2006, pp. 158–159) when given two different copulas and respective parameters as arguments. The function concordCOP just dispatches to tauCOP. A useful relation is

$$\iint_{\mathcal{I}^2} \mathbf{C}_1(u, v) d\mathbf{C}_2(u, v) = \frac{1}{2} - \iint_{\mathcal{I}^2} \frac{\delta}{\delta u} \mathbf{C}_1(u, v) \frac{\delta}{\delta v} \mathbf{C}_2(u, v) du dv,$$

where $\mathbf{C}_1(u, v)$ is the first copula and $\mathbf{C}_2(u, v)$ is the second copula.

Nelsen *et al.* (2001, p. 281) lists several measures of association defined by the concordance function:

1. $\tau_{\mathbf{C}} = \mathcal{Q}(\mathbf{C}, \mathbf{C})$: (Kendall Tau; tauCOP);
2. $\rho_{\mathbf{C}} = 3 \cdot \mathcal{Q}(\mathbf{C}, \mathbf{\Pi})$: (Spearman Rho; rhoCOP);
3. $\gamma_{\mathbf{C}} = 2 \cdot \mathcal{Q}(\mathbf{C}, [\mathbf{M} + \mathbf{W}]/2)$: (Gini Gamma; giniCOP); and
4. $\psi_{\mathbf{C}} = \frac{3}{2} \cdot \mathcal{Q}(\mathbf{C}, \mathbf{M}) - \frac{1}{2}$: (Spearman Footrule; footCOP).

Author(s)

W.H. Asquith

References

- Durante, F., and Sempi, C., 2015, Principles of copula theory: Boca Raton, CRC Press, 315 p.
- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Nelsen, R.B., Quesada-Molina, J.J., Rodríguez-Lallena, J.A., and Úbeda-Flores, M., 2001, Distribution functions of copulas—A class of bivariate probability integral transforms: Statistics and Probability Letters, v. 54, no. 3, pp. 277–282.

See Also

blomCOP, footCOP, giniCOP, hoefCOP, rhoCOP, wolfCOP, joeskewCOP, uvlmoms, derCOP, derCOP2, kfuncCOP

Examples

```
## Not run:
tauCOP(cop=PSP) # 1/3
# Now compute Kendall Tau via integration of the Kendall Function.
# 3 - 4*integrate(function(t) kfuncCOP(t, cop=PSP), 0, 1)$value # 0.3333314
## End(Not run)

## Not run:
tauCOP(cop=PSP, brute=TRUE) # 0.3306625
# CPU heavy example showing that the dual-integration (fast) results in
# a Kendall Tau that matches a sample version
dotau <- function(n) {
  uv <- simCOP(n=n, cop=PSP, ploton=FALSE, points=FALSE)
  return(cor(uv$U, uv$V, method="kendall"))
}
```

```

set.seed(817600)
taus <- replicate(100, dotau(100))
tau.sample <- mean(taus); print(tau.sample) # 0.3342034
## End(Not run)

## Not run:
# Nelsen (2006, pp. 160-161, numeric results shown therein)
# The rational values or integers may be derived analytically.
tauCOP(cop=M, cop2=M) # 1, correct
tauCOP(cop=M, cop2=P) # 1/3, correct
tauCOP(cop=P, cop2=M) # 1/3, correct
tauCOP(cop=M, cop2=W) # 0, correct
tauCOP(cop=W, cop2=M) # throws warning, swaps copulas, == tauCOP(M,W)
tauCOP(cop=W, cop2=P) # throws warning, swaps copulas, approx. -1/3
tauCOP(cop=P, cop2=W) # -1/3, correct
tauCOP(cop=P, cop2=P) # 0, correct
tauCOP(cop=M, cop2=W, brute=TRUE) # 0, correct
## End(Not run)

## Not run:
para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
             para1=0.00395, para2=4.67, alpha=0.9392, beta=0.5699)
tauCOP(cop=composite2COP, para=para) # -0.4671213

para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
             para1=0.14147, para2=20.96, alpha=0.0411, beta=0.6873)
tauCOP(cop=composite2COP, para=para) # +0.1950727

para <- list(cop1=PLACKETTcop, cop2=PLACKETTcop,
             para1=0.10137, para2=4492.87, alpha=0.0063, beta=0.0167)
# Theoretical attempt fails because para2 is large and thus a singularity
# is emerging and internal copula swapping does not help.
tauCOP(cop=composite2COP, para=para) # fails (0.94+- .01)
tauCOP(cop=composite2COP, para=para, brute=TRUE) # about 0.94+- .01
## End(Not run)

```

Tcop

The *t*-Student Copula

Description

The *t*-Student copula (Salvadori *et al.*, 2007, pp. 255–256) is

$$\mathbf{C}_{\Theta, \nu}(u, v) = \mathbf{T}(u, v; \Theta, \nu) = \int_{-\infty}^{t_{\nu}^{(-1)}(u)} \int_{-\infty}^{t_{\nu}^{(-1)}(v)} \frac{1}{2\pi\sqrt{1-\Theta^2}} \exp\left(-\frac{s^2 - 2\Theta st + t^2}{\nu(1-\Theta^2)}\right)^{-(\nu+2)/2} ds dt,$$

where $\Theta \in [-1, 1]$, $\nu \geq 1$ (integer), and $t_{\nu}^{(-1)}(x)$ is the quantile function of the *univariate t-distribution*. The copula, as $\Theta \rightarrow -1^+$, limits to the *countermonotonicity copula* ($\mathbf{W}(u, v)$; **W**), as

$\Theta \rightarrow 0$ limits, to the *independence coupla* (**P**(u, v) if ν becomes large; **P**), and as $\Theta \rightarrow 1^-$, limits to the *comonotonicity copula* (**M**(u, v); **M**). The copula has *lower-tail dependency* and *upper-tail dependency parameters* that are nonzero if $\Theta > 0$ and both parameters equal to

$$\lambda_{L|U} = 2t_{\nu+1} \left(-\frac{\sqrt{\nu+1}\sqrt{1-\Theta}}{\sqrt{1+\Theta}} \right),$$

which tend to zero as $\nu \rightarrow \infty$, which are those for the *Normal copula* (**NORMcop**). The *Spearman Rho* (**rhoCOP**) is $\rho_C = (6/\pi) \cdot \text{asin}(\Theta/2)$ and *Kendall Tau* (**tauCOP**) is $\tau_C = (2/\pi) \cdot \text{asin}(\Theta)$ and are the same as for **NORMcop**. The parameter Θ is readily computed by $\Theta = 2 \cdot \sin(\pi \cdot \rho_C/6)$ or by $\Theta = \sin(\pi \cdot \tau_C/2)$.

Usage

```
Tcop(u, v, para=NULL, rho=NULL, tau=NULL, taildep=NULL, fit=c("rho", "tau"), ...)
```

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction;
para	A vector (two element) of parameters— Θ and ν parameters of the copula and note that ν will be silently cast as an integer after rounding to zero digits internally for <code>mvtnorm::pmvt()</code> ;
rho	Optional Spearman Rho from which the parameter will be estimated and presence of rho trumps tau;
tau	Optional Kendall Tau from which the parameter will be estimated;
taildep	Optional lower/upper tail dependency coefficient to try to fit the parameters for the give rho or tau;
fit	If para, rho, and tau are all NULL, then the u and v represent the sample. The measure of association by the fit declaration will be computed and the parameter estimated subsequently. The fit has no other utility than to trigger which measure of association is computed internally by the cor function in R; and
...	Additional arguments to pass.

Value

Value(s) for the copula are returned. Otherwise if either rho or tau is given, then the Θ and ν are computed and a list having

para	The parameters Θ and ν (if computable, refer to message) and note that ν will be silently cast as an integer after rounding to zero digits internally for <code>mvtnorm::pmvt()</code> ;
rho	Spearman Rho if the rho is given; and
tau	Kendall Tau if the tau is given but also if both rho and tau are NULL as mentioned next.
lambdaUL	Computed tail dependency (if computable);

```

message      Helpful message;
lambdaUL_if_nu_were_eqone
              Computed tail dependency if  $\nu$  were to equal 1;

```

and if `para=NULL` and `rho` and `tau=NULL`, then the values within `u` and `v` are used to compute Spearman Rho (`fit="rho"`) or Kendall Tau (`fit="tau"`) and then compute the Θ and then attempt to compute the ν from the `taildep`, and these are returned in the aforementioned list.

Note

A mimic of the wrappers on `mvtnorm::pmvt()` from the **copula** package (v1.1-6) are made within `Tcop`. The rest of the implementation is unique to **copBasic**. An implementation of a function in the **copBasic** style that would natively interconnect to **copula** package version of the copula could be as follows:

```

"Tcop" <-      # pCoupla() from package copula is analogous to COP()
function(u,v, para=NULL) {
  if(length(u) == 1) u <- rep(u, length(v)) # see asCOP() for reasoning of
  if(length(v) == 1) v <- rep(v, length(u)) # this "vectorization" hack
  para <- copula::tCopula(c(para[1]), dim=2, df=as.integer(para[2]))
  return( copula::pCopula(matrix(c(u,v), ncol=2), para) )
}

```

Author(s)

W.H. Asquith

References

Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, *Extremes in Nature—An approach using copulas*: Springer, 289 p.

See Also

[NORMcop](#)

Examples

```

## Not run:
para <- Tcop(para=NULL, rho=0.9, taildep=0.3) # 0.907981 22.000000
UV <- simCOP(1000, cop=Tcop, para=para$para) # see the tail dependency
## End(Not run)

```

tEVcop

The *t*-EV (Extreme Value) Copula**Description**

The *t*-EV copula (Joe, 2014, p. 189) is a limiting form of the *t*-Student copula (multivariate *t*-distribution) ([TCop](#)):

$$\mathbf{C}_{\rho,\nu}(u, v) = \mathbf{tEV}(u, v; \rho, \nu) = \exp(-(x + y) \times B(x/(x + y); \rho, \nu)),$$

where $x = -\log(u)$, $y = -\log(v)$, and letting $\eta = \sqrt{(\nu + 1)/(1 - \rho^2)}$ define

$$B(w; \rho, \nu) = w \times t_{\nu+1}(\eta[(w/[1 - w])^{1/\nu} - \rho]) + (1 - w) \times t_{\nu+1}(\eta[(1 - w)/w]^{1/\nu} - \rho),$$

where $t_{\nu+1}$ is the cumulative distribution function of the *univariate t-distribution* with $\nu + 1$ degrees of freedom. As $\nu \rightarrow \infty$, the copula weakly converges to the *Hüsler–Reiss copula* ([HRCop](#)) because the *t*-distribution converges to the normal (refer to **Examples** for a study of this copula).

The $\mathbf{tEV}(u, v; \rho, \nu)$ copula is a two-parameter option when working with extreme-value copula. There is a caveat though. Demarta and McNeil (2004) conclude that “the parameter of the *Gumbel–Hougaard* ([GHcop](#)) or *Galambos* ([GLcop](#)) A-functions [the *Pickend dependence function* and B-function by association] can always be chosen so that the curve is extremely close to that of the *t*-EV A-function for any values of ν and ρ . The implication is that in all situations where the *t*-EV copula might be deemed an appropriate model then the practitioner can work instead with the simpler Gumbel or Galambos copulas.”

Usage

```
tEVcop(u, v, para=NULL, ...)
```

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction;
para	A vector (two element) of parameters in ρ and ν order; and
...	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Note

Note, Joe (2014) shows $x = \log(u)$ (note absence of the minus sign)—this is not correct.

Author(s)

W.H. Asquith

References

- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
- Demarta, S., and McNeil, A.J., 2004, The t copula and related copulas: International Statistical Review, v. 33, no. 1, pp. 111–129, doi:10.1111/j.17515823.2005.tb00254.x

See Also

[GHcop](#), [GLcop](#), [HRcop](#), [Tcop](#)

Examples

```
## Not run:
tau <- 1/3 # Example from copula::evCopula.Rd
tev.cop <- copula::tevCopula(copula::iTau(copula::tevCopula(), tau))
copula::pCopula(c(0.1,.5), copula=tev.cop) # 0.07811367
tEvcop(0.1, 0.5, para=slot(tev.cop, "parameters")) # 0.07811367
## End(Not run)

## Not run:
nsim <- 2000; pargh <- c(5, 0.5, 0.5)
UV <- simCOP(nsim, cop=GHcop, para=pargh)
U <- lmomco::pp(UV[,1], sort=FALSE)
V <- lmomco::pp(UV[,2], sort=FALSE)
RT <- mleCOP(u=U, v=V, cop=tEvcop, init.para=c(0.5, log(4)),
             parafn=function(k) return( c(k[1], exp(k[2])) ) )
partev <- RT$para

FT <- simCOP(nsim, cop=tEvcop, para=RT$para)

tauCOP(cop=GHcop, para=pargh )
tauCOP(cop=tEvcop, para=partev)

tauCOP(cop=GHcop, para=pargh ) # [1] 0.3003678
tauCOP(cop=tEvcop, para=partev) # [1] 0.3178904

densityCOPplot(cop=GHcop, para=pargh)
densityCOPplot(cop=tEvcop, para=partev, ploton=FALSE, contour.col="red") #
## End(Not run)

## Not run:
# A demonstration Joe (2014, p. 190) for which tEvcop() has
# upper tail dependence parameter as
para <- c(0.8, 10)
lamU <- 2 * pt( -sqrt( (para[2]+1) * (1-para[1]) / (1+para[1]) ), para[2]+1)
"tEvcop.copula" <- function(u,v, para=NULL, ...) {
  if(length(u) == 1) u <- rep(u,length(v)); if(length(v)==1) v <- rep(v,length(u))
  return(copula::pCopula(matrix(c(u,v), ncol=2),
                             copula::tevCopula(param=para[1], df=para[2])))
}
lamU.copBasic <- taildepCOP(cop=tEvcop, para)$lambdaU
lamU.copula <- taildepCOP(cop=tEvcop.copula, para)$lambdaU
```

```

print(c(lamU, lamU.copBasic, lamU.copula))
#[1] 0.2925185 0.2925200 0.2925200 # So, we see that they all match.
## End(Not run)

## Not run:
# Convergence of tEvcop to HRcop as nu goes to infinity.
nu <- 10^(seq(-4, 2, by=0.1)) # nu right end point rho dependent
rho <- 0.7 # otherwise, expect to see 'zeros' errors on the plot()
# Compute Blomqvist Beta (fast computation is reason for choice)
btEV <- sapply(nu, function(n) blomCOP(tEvcop, para=c(rho, n)))
limit.thetas <- sqrt(2 / (nu*(1-rho))) # for nu --> infinity HRcop
thetas <- sapply(btEV, function(b) {
  uniroot(function(l, blom=NA) { blom - blomCOP(HRcop, para=l) },
    interval=c(0,10), blom=b)$root })
plot(limit.thetas, thetas, log="xy", type="b",
  xlab="Theta of HRcop via limit nu --> infinity",
  ylab="Theta from Blomqvist Beta equivalent HRcop to tEvcop")
abline(0,1)
mtext(paste0("Notice the 'weak' convergence to lower left, and \n",
  "convergence increasing with rho"))
# Another reference of note
# https://mediatum.ub.tum.de/doc/1145695/1145695.pdf (p.39) #
## End(Not run)

```

Description

The *Triangular copula* (Nelsen, 2006, exam. 3.3, pp. 59–60; exer. 3.7, pp. 64–65) is

$$\mathbf{C}_{\Theta}(u, v) = \mathbf{TRI}(u, v)$$

for $\Theta \in (0, -1)$. The copula, as $\Theta \rightarrow 0$ limits, to the *countermonotonicity coupla* ($\mathbf{W}(u, v)$; $\mathbf{W}$), as $\Theta \rightarrow 1$ limits to the *comonotonicity copula* ($\mathbf{M}(u, v)$; $\mathbf{M}$). The parameter Θ is readily computed from a *Spearman Rho* ($\mathbf{rhoCOP}$) or *Kendall Tau* ($\mathbf{tauCOP}$) by $\rho_{\mathbf{C}} = \tau_{\mathbf{C}} = 2\Theta - 1$. However in general for finite samples, the sample versions of Spearman Rho and Kendall Tau will not be equal and different Θ wil result.

Usage

```
TRIcop(u, v, para=NULL, rhotau=NULL, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction;
<code>para</code>	A vector (single element) of parameters—the Θ parameter of the copula;
<code>rhotau</code>	Optional Spearman Rho or Kendall Tau; and
<code>...</code>	Additional arguments to pass.

Value

Value(s) for the copula are returned. Otherwise if rhotau is given, then the Θ is computed and a list having

para The parameter Θ , and
 rhotau Spearman Rho or Kendall Tau.

and if para=NULL and rhotau=NULL, then the values within u and v are used to compute Spearman Rho or Kendall Tau and then compute the parameter, and these are returned in the aforementioned list.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[M](#), [W](#)

Examples

```
# Concerning Nelsen (2006, exer. 5.6, p. 171)
rhoCOP(cop=TRICop, para=0.25) # -0.4999916
tauCOP(cop=TRICop, para=0.25) # -0.5

## Not run:
for(i in seq_len(4)) {
  para <- list(cop=TRICop, para=0.25, reflect=i)
  uv <- simCOP(100, cop=COP, para=para, seed=1, col=i, ploton=! as.logical(i-1))
} #
## End(Not run)
```

uvlmoms

Bivariate Skewness after Joe (2014) or the Univariate L-moments of Combined U and V

Description

Joe (2014, pp. 65–66) suggests two quantile-based measures of *bivariate skewness* defined for uniform random variables U and V combined as either $\psi_{u+v-1} = u + v - 1$ or $\psi_{u-v} = u - v$ for which the $E[u] = E[v] = 0$. The bivariate skewness is the quantity η :

$$\eta(p; \psi) = \frac{x(1-p) - 2x(\frac{1}{2}) + x(p)}{x(1-p) - x(p)},$$

where $0 < p < \frac{1}{2}$, $x(F)$ is the quantile function for nonexceedance probability F for either the quantities $X = \psi_{u+v-1}$ or $X = \psi_{u-v}$ using either the empirical quantile function or a fitted distribution. Joe (2014, p. 66) reports that $p = 0.05$ to “achieve some sensitivity to the tails.” How these might be related (intuitively) to L-coskew (see function `lcomoms2()` of the **lmomco** package) of the L-comoments or bivariate L-moments (**bilmom**s) is unknown, but see the **Examples** section of **joeskewCOP**.

Structurally the above definition for η based on quantiles is oft shown in comparative literature concerning L-moments. But why stop there? Why not compute the L-moments themselves to arbitrary order for η by either definition (the `uvlmoms` variation)? Why not fit a distribution to the computed L-moments for estimation of $x(F)$? Or simply compute “skewness” according to the definition above (the `uvskew` variation).

Usage

```
uvlmoms(u,v=NULL, umv=TRUE, p=NA,   type="gno", getlmoms=TRUE, ...)

uvskew( u,v=NULL, umv=TRUE, p=0.05, type=6,      getlmoms=FALSE, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction and if <code>NULL</code> then <code>u</code> is treated as a two column <code>R</code> data frame;
<code>umv</code>	A logical controlling the computation of ψ : $\psi = u - v$ (<code>umv = TRUE</code>) or $\psi = u + v - 1$ (<code>umv = FALSE</code>). The “m” is to read “minus”;
<code>p</code>	A suggested p value is <code>p = 0.05</code> . If <code>is.na(NA)</code> , then <code>getlmoms</code> is set to <code>TRUE</code> (see below);
<code>type</code>	The <code>type</code> argument is mutable, and is a syntax match to the canonical use in package lmomco . Variation from that package however is permitted. Either <code>type</code> is an integer between 1 and 9 selecting one of the nine quantile algorithms described for the quantile function in <code>R</code> . The default 6 uses the <i>Weibull plotting positions</i> and differs from the <code>R</code> default of 7. Otherwise <code>type</code> must be a valid distribution abbreviation for the lmomco package as in the abbreviation list <code>dist.list</code> function of that package. The <code>gno</code> shown as a default for the generalized normal distribution (see distribution type “ <code>gno</code> ” in package lmomco);
<code>getlmoms</code>	A logical triggering whether the L-moments of either ψ_{u+v-1} or ψ_{u-v} are returned instead computing the above definition of “skewness;” and
<code>...</code>	Additional arguments to pass to the lmomco function <code>lmoms</code> , such as the number of L-moments <code>nmoms</code> .

Value

An `R` list of the univariate L-moments of η is returned (see documentation for `lmoms` in the **lmomco** package). Or the skewness of η can be either (1) based on the empirical distribution based on plotting positions by the quantile function in `R` using the `type` as described, or (2) based on the fitted quantile function for the parameters of a distribution for the **lmomco** package.

Author(s)

W.H. Asquith

References

Asquith, W.H., 2011, Distributional analysis with L-moment statistics using the R environment for statistical computing: Createspace Independent Publishing Platform, ISBN 978–146350841–8.

Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.

See Also[COP](#)**Examples**

```
## Not run:
set.seed(234)
UV <- simCOP(n=100, cop=GHCop, para=1.5, graphics=FALSE)
lmr <- uvlmoms(UV); print(lmr) # L-kurtosis = 0.16568268
uvskew(UV, p=0.10)           # -0.1271723
uvskew(UV, p=0.10, type="gno") # -0.1467011
## End(Not run)

## Not run:
pss <- seq(0.01,0.49, by=0.01)
ETA <- sapply(1:length(pss), function(i) uvskew(UV, p=pss[i], type=5, uvm1=FALSE) )
plot(pss, ETA, type="l", xlab="P FACTOR", ylab="BIVARIATE SKEWNESS") #
## End(Not run)
```

vuongCOP

*The Vuong Procedure for Parametric Copula Comparison***Description**

Perform the *Vuong Procedure* following Joe (2014, pp. 257–258). Consider two copula densities $f_1 = c_1(u, v; \Theta_1)$ and $f_2 = c_2(u, v; \Theta_2)$ for two different bivariate copulas $C_1(\Theta_1)$ and $C_2(\Theta_2)$ having respective parameters Θ_1 and Θ_2 that provide the “closest” *Kullback–Leibler Divergence* from the true copula density $g(u, v)$.

The difference of the Kullback–Leibler Divergence ([kullCOP](#)) of the two copulas from the true copula density can be measured for a sample of size n and bivariate sample realizations $\{u_i, v_i\}$ by

$$\hat{D}_{12} = n^{-1} \sum_{i=1}^n D_i,$$

where $\hat{D}_{12}$ is referred to in the **copBasic** package as the “Vuong D ” and D_i is defined as

$$D_i = \log \left[\frac{f_1(u_i, v_i; \Theta_2)}{f_2(u_i, v_i; \Theta_1)} \right].$$

The variance of $\hat{D}_{12}$ can be estimated by

$$\hat{\sigma}_{12}^2 = (n-1)^{-1} \sum_{i=1}^n (D_i - \hat{D}_{12})^2.$$

The sample estimate and variance are readily turned into the $100 \times (1 - \alpha)$ confidence interval by

$$\hat{D}_{12}^{(lo)} < \hat{D}_{12} < \hat{D}_{12}^{(hi)},$$

where, using the quantile (inverse) function of the t-distribution $\sim \mathcal{T}^{(-1)}(F; df=(n-2))$ for nonexceedance probability F and $n-2$ degrees of freedom for n being the sample size, the confidence interval is

$$\hat{D}_{12} - \mathcal{T}^{(-1)}(1 - \alpha/2) \times \hat{\sigma}_{12} / \sqrt{n} < \hat{D}_{12} < \hat{D}_{12} + \mathcal{T}^{(-1)}(1 - \alpha/2) \times \hat{\sigma}_{12} / \sqrt{n}.$$

Joe (2014, p. 258) reports other interval forms based (1) on the Akaike (AIC) correction and (2) on the Schwarz (BIC) correction, which are defined for AIC as

$$\text{AIC} = \hat{D}_{12} - (2n)^{-1} \log(n) \left[\dim(\Theta_2) - \dim(\Theta_1) \right] \pm \mathcal{T}^{(-1)}(1 - \alpha/2) \times \hat{\sigma}_{12} / \sqrt{n},$$

and for BIC as

$$\text{BIC} = \hat{D}_{12} - (2n)^{-1} \log(n) \left[\dim(\Theta_2) - \dim(\Theta_1) \right] \pm \mathcal{T}^{(-1)}(1 - \alpha/2) \times \hat{\sigma}_{12} / \sqrt{n}.$$

The AIC and BIC corrections incorporate the number of parameters in the copula and for all else being equal the copula with the fewer parameters is preferable. If the two copulas being compared have equal number of parameters then the AIC and BIC equate to $\hat{D}_{12}$ and the same confidence interval because the difference $[\dim(\Theta_2) - \dim(\Theta_1)]$ is zero.

Joe (2014, p. 258) reports that these three intervals can be used for *diagnostic inference* as follows. If an interval contains 0 (zero), then copulas $C_1(\Theta_1)$ and $C_2(\Theta_2)$ are not considered significantly different. If the interval does not contain 0, then copula $C_1(\Theta_1)$ or $C_2(\Theta_2)$ is the better fit depending on whether the interval is completely below 0 (thus $C_1(\Theta_1)$ better fit) or above 0 (thus $C_2(\Theta_2)$ better fit), respectively. Joe (2014) goes on to emphasize that “the procedure compares different [copulas] and assesses whether they provide similar fits to the data. [The procedure] does not assess whether [either copula] is a good enough fit.”

Usage

```
vuongCOP(u, v=NULL, cop1=NULL, cop2=NULL, para1=NULL, para2=NULL,
         alpha=0.05, method=c("D12", "AIC", "BIC"),
         the.zero=.Machine$double.eps^0.25, ...)
```

Arguments

<code>u</code>	Nonexceedance probability u in the X direction;
<code>v</code>	Nonexceedance probability v in the Y direction and if NULL then <code>u</code> is treated as a two column R data frame;
<code>cop1</code>	A copula function corresponding to copula f_1 in the Vuong Procedure;

para1	Vector of parameters or other data structure, if needed, to pass to the copula f_1 ;
cop2	A copula function corresponding to copula f_2 in the the Vuong Procedure;
para2	Vector of parameters or other data structure, if needed, to pass to the copula f_2 ;
alpha	The α in the Vuong Procedure, which results in the $100 \times (1 - \alpha)$ confidence interval (two sided);
method	The interval to evaluate as to position of the respective statistic form the comparison of the two copulas;
the.zero	The value for “the zero” of the copula density function. This argument is the argument of the same name for densityCOP . The default here is intended to suggest that a tiny nonzero value for density will trap the numerical zero densities; and
...	Additional arguments to pass to the densityCOP function.

Value

An R list is returned having the following components:

title	A descriptive title of the procedure;
method	A textual description of the method setting;
result.text	A textual description of the result of the Vuong Procedure;
result	A value 1 if $C_1(\Theta_1)$ is better fit, 2 if copula $C_2(\Theta_2)$ is better fit, and 0 if neither is better ($\hat{D}_{12} = 0$), and NA including the likely(?) erroneous situation of $C_1(\Theta_1) \equiv C_2(\Theta_2)$;
p.value	The two-sided p-values of the Vuong Procedure inclusive of AIC and BIC;
D12	A named vector of the lower and upper bounds of Vuong D at the respective confidence interval percentage along with $\hat{D}_{12}$ and σ_{12}^2 ;
AIC	A named vector of the lower and upper bounds of Vuong AIC at the respective confidence interval percentage;
BIC	A named vector of the lower and upper bounds of Vuong BIC at the respective confidence interval percentage; and
parameters	A named vector of the alpha, sample size, value for the t-distribution quantile $qt(1-\alpha/2, df=n)$, and $\hat{\sigma}_{12}$.

Note

The [vuongCOP](#) function along with [kullCOP](#) and features of function [densityCOPplot](#) represent collective milestones towards *copula inference* and diagnostics post fitting of copulas to the usual measures of association such as the *Kendall Tau* (τ_K) and *Spearman Rho* (ρ_S) and their copula counterparts τ_C ([tauCOP](#)) and ρ_C ([rhoCOP](#)).

For an example application, imagine a problem of say low springflow risk at “nearby springs” that jointly should converge in the lower tail because drought usually has a strong regional impact. First, it is necessary to form a reflection of the *Gumbel–Hougaard copula* ($\mathbf{GH}(u, v; \Theta_{\mathbf{GH}})$; [GHcop](#)) but parameter estimation using τ_C is the same because sample $\hat{\tau}_K$ is invariant to reflection.

```
"rGHcop" <- function(u,v, ...) { u + v - 1 + GHcop(1-u, 1-v, ...) }
set.seed(385) # setting so that reported quantities here are reproducible
```

The prior code also sets a seed on the pseudo-random number generator so that reported values here are reproducible. The reflected $\mathbf{GH}(u, v; \Theta_{\mathbf{GH}})$ is denoted $\mathbf{rGH}(u, v; \Theta_{\mathbf{rGH}})$.

Second, the $\mathbf{PSP}(u, v)$ copula (**PSP**) is chosen as the parent distribution, and this copula has no parameter. The **PSP** has lower-tail dependency, which will be important as discussion unfolds. The following two lines of code establish a sample size to be drawn from the **PSP** and then simulates a sample from that copula. The color grey is used for the simulated values on the figure produced by **simCOP**, which forms a background example of the joint structure of the **PSP** copula.

```
n <- 390
UV <- simCOP(cop=PSP, n=n, col=8, pch=16) # simulate and form the base image
```

By inspection of the so-produced graphic, it is obvious that there is contraction in the lower-left corner of the plot, which is a geometric representation of tail dependency. The lower-tail dependency thus phenomenologically says that there is joint interconnect during low springflow conditions—both springs are likely to be at low flow simultaneously. The variable UV contains the bivariate data as uniform variables (nonexceedance probabilities u and v).

The *Plackett copula* ($\mathbf{PL}(u, v; \Theta_{\mathbf{PL}})$; **PLACKETTcop**) and the $\mathbf{rGH}(u, v; \Theta_{\mathbf{rGH}})$ copula are chosen as candidate models of the “unknown” parent. Both **PL** and **rGH** copulas use different “measures of association” for their parameter estimation. Next, sample estimates of the copula parameters using *Schweizer and Wolff Sigma* $\hat{\sigma}_C$. The sample value computations and parameter estimates also are set as shown in the following code:

```
Wolf <- wolfCOP(para=UV, as.sample=TRUE) # 0.496943
paraPL <- uniroot(function(p)
  Wolf - wolfCOP(cop=PLACKETTcop, para=p), c(1,30))$root
paraGH <- uniroot(function(p)
  Wolf - wolfCOP(cop=rGHcop, para=p), c(1,30))$root
```

STEP 1—Compute Kullback–Leibler sample size: The Kullback–Leibler Divergences ($\text{KL}(f|g)$ and $\text{KL}(g|f)$) are computed (**kullCOP**) for the evaluation of the sample size as appropriate for distinguishing between the two candidate copulas 95 percent of the time. The Kullback–Leibler sample size (n_{fg}) also is computed as the following code illustrates and provides additional commentary.

```
KL <- replicate(20, kullCOP(cop1=PLcop, para1=paraPL, # CPU intensive
  cop2=rGHcop, para2=paraGH, n=1E5)$KL.sample.size)
print(round(mean(KL))) # n_{fg} = 221 sample size
print(range(KL)) # 204 <-- n_{fg} --> 252 sample size range
```

Depending on the sample $\hat{\sigma}_C$ coming from the simulation of the parent **PSP** copula, the call to **kullCOP** will likely report different n_{fg} values because $n_{fg}(\mathbf{C}_1(\Theta_1), \mathbf{C}_1(\Theta_1))$. These sample sizes have a range for 20 replications of about $n_{fg} = 204\text{--}252$. The result here is $n_{fg} = 221$ and thus **the sample size $n = 390$ should be more than large enough to generally distinguish between the PL and rGH copulas at the respective sample measure of association.**

STEP 2—Perform the Vuong Procedure: The Vuong Procedure can now be completed. Now watch the copula and parameter order in the next code for mistakes, the author has purposefully switched

order here to draw attention to the need to make sure argument `cop1` has the correct parameter(s) for copula 1 (the **PL**). The two calls to `simCOP` are made to graphically superimpose these simulations on top of the parent **PSP**.

```
VD <- vuongCOP(UV, cop2=rGHcop, para2=paraGH, cop1=PLcop, para1=paraPL)
print(VD) # "Copula 2 better" or rGHcop (Gumbel-Hougaard is better)
set.seed(385) # seems harmless enough to reuse the seed to emphasize "fit"
TMP <-simCOP(cop=PLcop, para=paraPL,n=n,plot=FALSE,col="red", pch=16,cex=0.5)
set.seed(385) # seems harmless enough to reuse the seed to emphasize "fit"
TMP <-simCOP(cop=rGHcop,para=paraGH,n=n,plot=FALSE,col="green",pch=16,cex=0.5)
rm(TMP) # just cleaning up the workspace.
```

Further discussion of the Vuong Procedure is informative. Simply speaking, the result is that **the rGH (copula 2) has better fit than PL (copula 1)**. The 95-percent confident limits from the procedure for $\hat{D}_{12} = 0.049$ with p-value 0.0012, $\hat{\sigma}_{12} = 0.297$, and $n = 390$ are $0.0194 < \hat{D}_{12} < 0.0786$. This interval does not contain zero and is greater than zero and therefore a conclusion may be drawn that copula 2 has the better fit.

STEP 3—Comparison of lower-tail dependency parameters: What does the tail dependency do for inference? This can be checked by computing the lower-tail dependency parameters (λ_G^L ; `taildepCOP`) in the code that follows for each of the three copulas and the empirical copula with acknowledgment that true sample estimators do not quite exist. Numeric focus need only be on the lower tail, but the four graphics are informative.

```
taildepCOP(cop=PSP,          plot=TRUE)$lambdaL # = 1/2
taildepCOP(cop=PLcop,      para=paraPL, plot=TRUE)$lambdaL # = ZERO
taildepCOP(cop=rGHcop,     para=paraGH, plot=TRUE)$lambdaL # = 0.429
taildepCOP(cop=EMPIRCop,   para=UV,      plot=TRUE)$lambdaL # = 0.328
```

The important aspect of the graphics by `taildepCOP` is that the **rGH** has lower-tail dependency whereas the **PL** does not. So, based on inspection **rGH** is superior given that we known **PSP** was the true parent. The empirical estimate of the $\hat{\lambda}_G^L = 0.328$ through the `EMPIRCop` copula indicates that its lower-tail dependency is closer to that of the **rGH** relative to **PL** and thus **quantitatively by lower-tail dependency the rGH has a superior fit**.

Therefore the **rGH** has a tail dependency more similar to the true model compared to the **PL**. Hence for this example, the **rGH** is clearly a superior fitting model in terms of the *Vuong Procedure* (fit alone) and the λ_G^L then is used as a follow up to shown that the **rGH** might be “good enough” an approximation to the **PSP**. The efficacy of reflecting the **GH** copula into a “new” form as **rGH** is demonstrated. Users are strongly encouraged to review the so-produced graphic from the `simCOP` call several listings back for $n = 390$, and lastly, this example is one for which absence of the argument `snv` (standard normal variate [scores]) by `simCOP` makes the tail dependency issue for the sample size more prominent in the graphic.

STEP 4—Qualitatively compare using copula density plots: Graphical depiction of copula density contours by the `densityCOPplot` function supports the conclusion that the **rGH** is the superior model relative to the **PL**. The so-produced graphic obviously shows that **the rGH strongly mimics the shape of the parent PSP**.

```
densityCOPplot(cop=PSP, contour.col=8) # grey is the parent bivariate density
densityCOPplot(cop=PLcop, para=paraPL, contour.col="green", ploton=FALSE)
densityCOPplot(cop=rGHcop, para=paraGH, contour.col="red", ploton=FALSE)
```

STEP 5—Compute L-comoments of the data via simulation and estimate the sampling distributions:

An open research problem is the what if any role that *L-comoments* might play in either copula estimation or inference. (There being very little literature on the topic?) Because a measure of association was used for parameter estimation, the L-correlation is uninformative, but a comparison is conceptually useful. The $\hat{\sigma}_C = 0.4969$ and *Spearman Rho* of the data $\hat{\rho}_S$ and the L-correlations $\hat{\rho}_S \approx \tau_2^{[12]} \approx \tau_2^{[21]} \approx 0.497$ are all similar as mandated by the mathematics.

Inference using L-coskew and L-cokurtosis seems possible. The following code listing is CPU intensive. First, the L-correlation, L-coskew, and L-cokurtosis values are computed from the simulated sample by the `lcomoms2()` function of the **lmomco** package. Second and third, the respective sampling distributions of these L-comoments (`lcomCOPpv`) for the two copulas are estimated.

```
UVlmr <- lmomco::lcomoms2(UV, nmom=4) # The sample L-comoments
# This execution will result in nonrejection of rGH copula.
GHlmr <- lcomCOPpv(n, UVlmr, cop=rGHcop, para=paraGH) # NONREJECTION
# LcomType      n      Mean  Lscale   Lskew   Lkurt sample.est p.value signif
#   Tau3[12] 390 -0.06952 0.01819 0.04505 0.12024 -0.11188 0.08795 .
#   Tau3[21] 390 -0.06739 0.02084 0.04104 0.12917 -0.10673 0.14162 -
#   Tau3[12:21] 390 -0.06845 0.01713 0.04930 0.11696 -0.10931 0.08161 .
#   Tau4[12] 390 0.04970 0.01682 -0.01635 0.10150 0.04183 0.38996 -
#   Tau4[21] 390 0.05129 0.01606 -0.06833 0.13798 0.07804 0.17470 -
#   Tau4[12:21] 390 0.05049 0.01329 -0.02045 0.12001 0.05994 0.35069 -

# This execution will result in rejection of Plackett copula.
PLlmr <- lcomCOPpv(n, UVlmr, cop=PLACKETTcop, para=paraPL) # REJECT PLACKETT
# LcomType      n      Mean  Lscale   Lskew   Lkurt sample.est p.value signif
#   Tau3[12] 390 -0.00267 0.02133 0.01556 0.09581 -0.11188 0.00129 **
#   Tau3[21] 390 -0.00112 0.02022 -0.00663 0.13338 -0.10673 0.00189 **
#   Tau3[12:21] 390 -0.00189 0.01757 0.00906 0.10226 -0.10931 0.00019 ***
#   Tau4[12] 390 0.00153 0.01652 -0.03320 0.12468 0.04183 0.07924 .
#   Tau4[21] 390 0.00361 0.01851 -0.01869 0.12052 0.07804 0.00929 **
#   Tau4[12:21] 390 0.00257 0.01362 -0.01194 0.10796 0.05994 0.00744 **
```

Because each copula was fit to a measure of association, the p-values for the L-correlations are all nonsignificant (noninformative because of how the copulas were fit), and therefore p-values quite near to the 50th percentile should be produced. So here, the L-correlation is noninformative on fit even though it might have some role because it is asymmetrical unlike that statistics τ_K and ρ_S . The results in variable `GHlmr` show no statistically significant entries (all p-values $> 0.05 = (\alpha = 0.1)/2$) for L-coskew and L-cokurtosis—the **rGH copula is not rejected**. The results in `PLlmr` show many p-values $< 0.05 = (\alpha = 0.1)/2$ for both L-coskew and L-cokurtosis—the **PL copula is rejected**. The experimental L-comoment inference shown is consistent with results with the Vuong Procedure.

The Vuong Procedure, however, does not address adequacy of fit—it just evaluates which copula fits better. The inspection of the lower tail dependency results previously shown ($\lambda_{PSP}^L = 1/2 \approx \lambda_{rGH}^U = 0.429$) along with the L-coskew and L-cokurtosis of the sample being well within the sample distribution suggests that the **rGH** is a adequate mimic of the **PSP** copula.

Some open research questions concern the numerical performance of the L-comoments as simulation sample size becomes large and whether or not the L-comoments should be computed on the probabilities $\{u, v\}$. Also should conversion to normal scores be made and if so, should adjustment

by the *Hazen plotting positions* ($u_i = (r_i - 0.5)/n$ for rank r_i) be made that Joe (2014) repeatedly advocates when standard normal variates (scores) [$z_i = \Phi^{(-1)}(u_i)$ for quantile function of standard normal distribution $\Phi(0, 1)$]? Collectively, Nelsen (2006) and Salvadori *et al.* (2007) are both silent on the matter of normal score conversion, and in conclusion Nelsen (2006), Salvadori *et al.* (2007), and Joe (2014) also are all silent on L-comoment applications with copulas.

Author(s)

W.H. Asquith

References

- Joe, H., 2014, Dependence modeling with copulas: Boca Raton, CRC Press, 462 p.
- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Salvadori, G., De Michele, C., Kottegoda, N.T., and Rosso, R., 2007, Extremes in Nature—An approach using copulas: Springer, 289 p.

See Also

[densityCOP](#), [kullCOP](#), [simCOP](#), [statTn](#), [mleCOP](#)

Examples

```
# See extended code listings and discussion in the Note section
# See Examples in mleCOP() (Last example therein might suggest a problem in the
# implied 95th percentile associated with n_fg above.
```

W

The Fréchet–Hoeffding Lower-Bound Copula

Description

Compute the *Fréchet–Hoeffding lower-bound copula* (Nelsen, 2006, p. 11), which is defined as

$$\mathbf{W}(u, v) = \max(u + v - 1, 0).$$

This is the copula of perfect anti-association (*countermonotonicity*, *perfectly negative dependence*) between U and V and is sometimes referred to as the *countermonotonicity copula*. Its opposite is the $\mathbf{M}(u, v)$ copula (*comonotonicity copula*; [M](#)), and statistical *independence* is the $\mathbf{\Pi}(u, v)$ copula ([P](#)).

Usage

`W(u, v, ...)`

Arguments

`u` Nonexceedance probability u in the X direction;
`v` Nonexceedance probability v in the Y direction; and
`...` Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[M](#), [P](#), [breveCOP](#), [kfuncCOP](#)

Examples

```
W(0.41, 0.60) # just barely touching the support, so small, 0.01
W(0.25, 0.45) # no contact with the support, so 0
W(1, 1) # total consumption of the support, so 1

## Not run:
# This example shows the impact of the "breve" permutation asymmetry addition
# to perfect negative correlation that though the plot of u,v changes with
# spread direction with sign of the breve, that the distribution function of
# the joint distribution still ranges uniformly 1 for breve = 0 down towards
# independence as breve approaches -1 and + 1. Then, repeat similarly for
# perfect positive correlation.
ff <- c(0.001, seq(0.01, 0.99, by=0.01), 0.999)
bs <- seq(-1, +1, by=0.1)
plot(c(0,1), c(0,1), type="n", xlab="Joint probability (Kendall function)",
     ylab="Nonexceedance probability of Kendall function")
for(b in bs) {
  lines(ff, kfuncCOP(ff, cop=breveCOP, para=list(cop=W, breve=b)),
        col=ifelse(b < 0, grey(0.8), "blue"), lwd=ifelse(b < 0, 6, 1))
}
# The top line will be W (breve = 0) and therefore its Kendall function is
# uniform distribution, and the lowest line is independence.
for(b in bs) {
  lines(ff, kfuncCOP(ff, cop=breveCOP, para=list(cop=M, breve=b)),
        col=ifelse(b < 0, grey(0.8), "seagreen"), lwd=ifelse(b < 0, 6, 1))
}
# The bottom line will be M (breve = 0) and therefore its Kendall function is
# perfect correlation. The top line by M and breve == +/-1 is independence
lines(ff, kfuncCOP(ff, cop=P), col="red", lwd=3) # draw independence in red
```

```

legend("bottomright", c("Breve in (0,+1] on M copula by breveCOP()",
                        "Breve in (0,+1] on W copula by breveCOP()",
                        "Perfect independence by P() copula",
                        "Breves in [-1, 0] in M or W copulas by breveCOP()"),
      lwd=c(1,1,3,6), col=c("seagreen", "blue", "red", "grey(0.8)")) #
## End(Not run)

```

wolfCOP

Schweizer and Wolff Sigma of a Copula

Description

Compute the measure of association known as *Schweizer–Wolff Sigma* σ_C of a copula according to Nelsen (2006, p. 209) by

$$\sigma_C = 12 \int \int_{T^2} |\mathbf{C}(u, v) - uv| \, du dv,$$

which is $0 \leq \sigma_C \leq 1$. It is obvious that this measure of association, without the positive sign restriction, is similar to the following form of *Spearman Rho* ([rhoCOP](#)) of a copula:

$$\rho_C = 12 \int \int_{T^2} [\mathbf{C}(u, v) - uv] \, du dv.$$

If a copula is *positively quadrant dependent* (PQD, see [isCOP.PQD](#)), then $\sigma_C = \rho_C$; conversely if a copula is *negatively quadrant dependent* (NQD), then $\sigma_C = -\rho_C$. However, a feature making σ_C especially attractive is that for random variables X and Y , which are not PQD or NQD—copulas that are neither larger nor smaller than Π —is that “ σ_C is often a better measure of [dependency] than ρ_C ” (Nelsen, 2006, p. 209).

Úbeda-Flores (2022) study the maximum difference between $\sigma_C - |\rho_C| \approx 0.60496$ for a special class of singular copula (see **Examples**), but states that it “remains an open problem to check if, for any copula $\mathbf{C}_\Theta$,” that the maximum listed here is the maximum difference possible.

Li *et al.* (2019) compare σ_C (they label it δ_i^E , “the extended delta index”) to alternative measures of dependency with emphasis on information content. Those authors also show that dependence can also be measured by

$$\delta_i = \frac{1}{2} \int \int_{T^2} |c(u, v) - 1| \, du dv,$$

where $c(u, v)$ is a *copula density* ([densityCOP](#)). Li *et al.* (2019, p. 45) “measure how much the model output Y and the input X_i are dependent,” and “importance of the inputs globally and quantitatively with model free and moment independent properties.”

By definition for two random variables U and V with cumulative distribution functions of $F(u)$ and $F(v)$, the variables are *independent*, if and only if, for all subsets (conditions) selecting values u and y of $U \leq u$ and $V \leq v$, when for all u and v , the combined cumulative distribution function is

$$H(x, y) = F(x)G(y) = \mathbf{C}(u, v) = uv.$$

Note, structural similarity between σ_C and the definition for [LzCOPpermsym](#). It seems to follow, by intuition, if therein, the copula on the left were to be replaced with the *permutation symmetric* (exchangeable symmetric) copula [isCOP.permsym](#), such as $\Pi = uv = vu$, then the copula on the right, now being, without loss of generality, $C(u, v)$ or $C(v, u)$, then σ_C must respond partly to dependence existing in the presence of *permutation asymmetry*.

Usage

```
wolfCOP(cop=NULL, para=NULL, as.sample=FALSE, brute=FALSE, delta=0.002,
        nlarge=500, usefastgrid=TRUE, ...)
```

Arguments

cop	A copula function;
para	Vector of parameters or other data structure, if needed, to pass to the copula;
as.sample	A logical controlling whether an optional R data.frame in para is used to compute the $\hat{\sigma}_C$ (see Details). If set to -1, then the message concerning CPU effort will be suppressed. This feature of controlling the message is not applicable if the fast computation by usefastgrid is set and sample size is larger than nlarge as the double summation in the Pózos <i>et al.</i> (2015) is bypassed in favor of EMPIRgrid_fast ;
brute	Should brute force be used instead of two nested integrate() functions in R to perform the double integration;
delta	The du and dv for the brute force (brute=TRUE) integration;
nlarge	A large sample size for which to consider the fast-grid computation if usefastgrid is true;
usefastgrid	A logical controlling whether the fast-grid computation is acceptable to the user; and
...	Additional arguments to pass to the copula itself, to EMPIRgrid_fast , or the integrate() function.

Details

A natural estimator for σ_C is through the *empirical copula* (Póczos *et al.*, 2015) and can be computed as

$$\hat{\sigma}_C = \frac{12}{n^2 - 1} \sum_{i=1}^n \sum_{j=1}^n \left| \hat{C}_n \left(\frac{i}{n}, \frac{j}{n} \right) - \frac{i}{n} \times \frac{j}{n} \right|,$$

where $\hat{C}_n$ is the simplest empirical copula of

$$\hat{C}_n \left(\frac{i}{n}, \frac{j}{n} \right) = \frac{1}{n} [\# \text{ of } (U_k \leq U_i, V_k \leq V_j)]$$

The sample version of $\hat{\sigma}_C$ is triggered by wolfCOP(para=D, as.sample=TRUE) for the double summation if usefastgrid is false and or usefastgrid is true and the same size is smaller than nlarge. A Monte Carlo integration version potentially suitable to very large sample sizes or difficult to integrate copula families is located at [wolfCOPsamc](#), but the function itself is likely of limited general

use because of the implementation of `EMPIRgrid_fast` inside `wolfCOP`. It is upon the fast estimator listed above that the extensive simulations leading to `wolfCOPtest` are based. In general, there are several asymptotically equivalent definitions of the empirical copula and some of which are implemented in `EMPIRcop`.

Consideration of speed performance between the `as.sample=TRUE` for the sample estimator argued by Póczos *et al.*, 2015) (refer to section **Note**), the `EMPIRgrid_fast` is measurably faster by about $n \geq 500$. The following code shows numerical equivalence between these estimators, but use of other *empirical copula* will have divergence unless samples are large:

```
"MYwolfCOPgrid" <- function(uv, ctype="1/n", ...) {
  ef <- EMPIRgrid_fast(para=uv, gridonly=FALSE, ctype=ctype)
  pf <- ef$empcop; n <- nrow(uv)
  gu <- as.numeric( rownames(ef$empcop) ) # gv <- gu (can compute w/o gv)
  for(i in seq_len(nrow(pf))) pf[i,] <- gu * gu[i]
  return(12 * sum(abs(ef$empcop - pf)) / (n^2 - 1))
}

ns <- as.integer( 10^seq(1, 3.3, by=0.1) ) # some sample sizes
WG <- WS <- vector(mode="numeric", length(ns))
for(i in seq_len(length(ns))) {
  uv <- simCOP(n=ns[i], cop=P, graphics=FALSE)
  WG[i] <- wolfCOP( para=uv, as.sample=TRUE, usefastgrid=FALSE)
  WS[i] <- MYwolfCOPgrid(uv)
}

opts <- par(xpd=NA, no.readonly=FALSE)
df <- data.frame(n=ns, wolf_grid=WG, wolf_sample=WS)
plot(df$n, df$wolf_sample, type="n", xlim=c(10, 10000), ylim=c(0, 0.60),
     xaxs="i", yaxs="i", log="x", las=1,
     xlab="Sample size of Independence copula",
     ylab="Schweizer-Wolff Sigma")
points(df$n, df$wolf_sample, pch=21, bg="white")
points(df$n, df$wolf_grid, pch=16, cex=0.6)
par(opts) #
```

Value

The value for σ_C is returned, and the sample $\hat{\sigma}_C$ computation versions have `names()` attached to verify to the user which subsystem was used given the `nlarge` and `usefastgrid` arguments.

Note

An extended example is informative. First, declare a composite of two different Plackett copulas (`PLcop`) and simulate a few hundred values. The parameter settings are such that the data will attain a look of independence.:

```
para <- list(cop1=PLcop, para1=0.145, alpha=0.81,
             cop2=PLcop, para2=21.90, beta=0.22)
D <- simCOP(n=300, cop=composite2COP, para=para,
```

```
cex=0.5, col=grey(0, 0.5), pch=16, seed=15)
```

Second, show that this copula is globally PQD (`isCOP.PQD`), but there is a substantial local NQD part of $\mathcal{L}^2$ space that clearly is NQD.

```
cols <- c("red3", "green4")
PQD <- isCOP.PQD(cop=composite2COP, para=para, uv=D)
message(PQD$global.PQD) # TRUE
points(D, col=cols[1+as.numeric(PQD$local.PQD)], lwd=2) # TRUE are green
```

This composited copula intersects, that is, passes through, the **P** copula. By the logic of Nelsen (2006), then the σ_C should be larger than ρ_C as shown below

```
wolfCOP(cop=composite2COP, para=para) # 0.08373378 (theoretical)
rhoCOP(cop=composite2COP, para=para) # 0.02845131 (theoretical)
hoefCOP(cop=composite2COP, para=para) # 0.08563823 (theoretical)
```

In fact, output above also shows Schweizer–Wolff Sigma to be larger than *Blomqvist Beta* (`blomCOP`), *Gini Gamma* (`giniCOP`), and *Kendall Tau* (`tauCOP`). Schweizer–Wolff Sigma captures the fact that although the symbols seems to plot randomly on the figure, symbol coloring for PQD and NQD clearly shows local dependency differences.

Author(s)

W.H. Asquith

References

- Li, L. Liu, Y., and Lu, Z., 2019, A new dependence measure for importance analysis—Application to an environmental model: Applied Mathematical Modelling, v. 74, pp. 43–61, [doi:10.1016/j.apm.2019.04.046](https://doi.org/10.1016/j.apm.2019.04.046).
- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.
- Póczos, B., Krishner, S., Pál, D., Szepesvári, C., and Schneider, J., 2015, Robust nonparametric copula based dependence estimators, accessed on August 11, 2015, at <https://www.cs.cmu.edu/~bapoczos/articles/poczos11nipscopula.pdf>.
- Schweizer, B., and Wolff, E.F., 1981, On nonparametric measures of dependence for random variables: The Annals of Statistics, v. 9, no. 4, pp. 879–855, <https://www.jstor.org/stable/2240856>.
- Úbeda-Flores, M., 2022, A note on the maximum difference between Schweizer and Wolff's Sigma and the absolute value of Spearman's Rho: Transactions on Fuzzy Sets and Systems, v. 1, no. 2, pp. 1–5, [doi:10.30495/tfss.2022.1956899.1024](https://doi.org/10.30495/tfss.2022.1956899.1024).

See Also

`wolfCOPsamc`, `wolfCOPtest`
`blomCOP`, `footCOP`, `giniCOP`, `hoefCOP`, `rhoCOP`, `tauCOP`, `joeskewCOP`, `uvlmoms`, `ORDSUMcop`

Examples

```
wolfCOP(cop=PSP) # 0.4784176

## Not run:
# Maximum Difference between Sigma and Absolute Spearman Rho
k <- 1/2 + sqrt(3)/6 # Ubeda-Flores (2022) or 1/2 - sqrt(3)/6
para <- list(cop=c(W, W), para=NULL, part=c(0, k, 1))
rho <- rhoCOP(ORDSUMcop, para=para)
wol <- wolfCOP(ORDSUMcop, para=para)
message("Wolf - |Rho| = ", wol - abs(rho))
# 0.604967382.... that is approx. 0.60496 given by Ubeda-Flores (2022).

ks <- seq(0.01, 0.99, by=0.002); rhos <- wols <- NULL
for(k in ks) {
  message(k, "-", appendLF=FALSE)
  para <- list(cop=c(W, W), para=NULL, part=c(0, k, 1))
  rhos <- c(rhos, abs( rhoCOP( cop=ORDSUMcop, para=para, subdivisions=200) ))
  wols <- c(wols,      wolfCOP(cop=ORDSUMcop, para=para, subdivisions=200) )
}
message("done")
plot( ks, rhos, type="l", lty=2, main="Ubeda-Flores (2022) Example 2.3",
      xlab="Theta of W,W shuffle", ylab="Value")
lines(ks, wols, lty=1); lines(ks, wols - rhos, lwd=3)
legend(0.3, 0.9, c("|Spearman Rho|", "Schweizer and Wolff Sigma",
                  "Schweizer and Wolff Sigma minus |Spearman Rho|"),
      bty="n", lty=c(2,1,1), lwd=c(1,1,2), cex=0.8)
abline(v=1/2 + c(-1, +1) * sqrt(3)/6, lty=4) # Ubeda-Flores (2022)
## End(Not run)

## Not run:
n <- 1000; UV <- simCOP(n=n, cop=N4212cop, para=7.53, graphics=FALSE, seed=1)
wolfCOP(cop=N4212cop, para=7.53) # 0.9884666 (theoretical)
wolfCOP(para=UV, as.sample=TRUE, usefastgrid=TRUE ) # 0.9873274 ( sample )
wolfCOP(para=UV, as.sample=TRUE, usefastgrid=FALSE) # 0.9873274 ( sample )
wolfCOPsamc(para=UV)$estimates[2] # 0.9861096 (Monte Carlo integration)
# Much faster as the sample size gets large to use the usefastgrid as shown
# in lieu of using the Monte Carlo. The wolfCOPsamc() was developed before
# the algorithm used for EMPIRgrid_fast and used by wolfCOP() was found.
## End(Not run)

## Not run:
# Redo D from Note section above
para <- list(cop1 =PLcop, cop2=PLcop,
             para1=0.145, para2=21.9, alpha=0.81, beta=0.22)
D <- simCOP(n=300, cop=composite2COP, para=para,
            cex=0.5, col=rgb(0, 0, 0, 0.2), pch=16)
PQD <- isCOP.PQD(cop=composite2COP, para=para, uv=D)
the.grid <- EMPIRgrid(para=D)
the.persp <- persp(the.grid$empcop, theta=-25, phi=20, shade=TRUE,
                  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")
empcop <- EMPIRcopdf(para=D) # data.frame of all points
points(trans3d(empcop$u, empcop$v, empcop$empcop, the.persp), cex=0.7,
```

```

col=rgb(0, 1-sqrt(empcop$empcop), 1, sqrt(empcop$empcop)), pch=16)
points(trans3d(empcop$u, empcop$v, empcop$empcop, the.persp),
       col=PQD$local.PQD+1, pch=1)

layout(matrix(c(1,2,3,4), 2, 2, byrow = TRUE), respect = TRUE)
PQD.NQD.cop <- gridCOP(cop=composite2COP, para=para)
Pi <- gridCOP(cop=P)
RHO <- PQD.NQD.cop - Pi; SIG <- abs(RHO)
the.persp <- persp(PQD.NQD.cop, theta=-25, phi=20, shade=TRUE, cex=0.5,
                  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")
mtext("The Copula that has local PQD and NQD", cex=0.5)
the.persp <- persp(Pi, theta=-25, phi=20, shade=TRUE, cex=0.5,
                  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")
mtext("Independence (Pi)", cex=0.5)
the.persp <- persp(RHO, theta=-25, phi=20, shade=TRUE, cex=0.5,
                  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")
mtext("Copula delta: Integrand of Spearman Rho", cex=0.5)
the.persp <- persp(SIG, theta=-25, phi=20, shade=TRUE, cex=0.5,
                  xlab="U VARIABLE", ylab="V VARIABLE", zlab="COPULA C(u,v)")
mtext("abs(Copula delta): Integrand of Schweizer-Wolff Sigma", cex=0.5) #
## End(Not run)

```

wolfCOPsamc

Monte Carlo Integration of Schweizer–Wolff Sigma

Description

Perform *Monte Carlo integration* for the measure of dependence known as *Schweizer–Wolff Sigma* $\sigma_C \in (0, 1)$ of an *empirical copula* ($C_n(u, v)$). This function can be useful as the `as.sample` computation provided by `wolfCOP` scales very poorly as sample size increases. Please refer to `wolfCOP` for more details, including theory, about σ_C . Sample estimation can be made by

$$\hat{\sigma}_C = 12 \int \int_{T^2} |C_n(u, v) - uv| \, du dv,$$

for which the integrals in the above formula are replaced with Monte Carlo integration provided by this function. Allowance is made internally to swap out $C_n(u, v)$ for a copula family $C(u, v)$, which means this function can be used for integration for families that might have numerical integration convergence problems in `wolfCOP`.

Usage

```

wolfCOPsamc(u=NULL, v=NULL, cop=EMPIRCOP, para=NULL, para_has_paras=FALSE,
            tol=0.005, minit=10, maxit=100, subdivisions=200, large_n=5E3,
            includeImoms=FALSE, trace=FALSE, verbose=FALSE, ...)

```

Arguments

<code>u</code>	Optional nonexceedance probability u in the X direction. This argument can also be a two-column data frame from which the <code>para</code> argument to EMPIRCop when <code>cop=EMPIRCop</code> is internally constructed;
<code>v</code>	Optional nonexceedance probability v in the Y direction from which the <code>para</code> argument to EMPIRCop when <code>cop=EMPIRCop</code> is internally constructed;
<code>cop</code>	The empirical copula, EMPIRCop , or a copula family;
<code>para</code>	Optional parameter content for the <code>para</code> argument to the specified copula in <code>cop</code> . For an empirical copula application (default), the <code>para</code> must be a two-column data frame, but if <code>cop</code> is set to a copula family, then this function can be used as an alternative σ_C computation using Monte Carlo integration in lieu of numerical integration exclusive to wolfCOP (refer to Examples how to use the function in this way);
<code>para_has_paras</code>	A logical to inform processing whether the <code>para</code> argument contains parameters for a copula family in lieu of <code>para</code> containing the sample for the empirical copula (<code>cop=EMPIRCop</code>). Internally, <code>para_has_paras</code> is used to disable random subsetting by setting the sample size variable to <code>large_n</code> . Technically, it is possible internally to detect whether <code>para_has_paras</code> needed to be true (even if false) as the number of random variates for the <code>halton()</code> function of the randtoolbox package is computed, and the code will silently proceed as if <code>para_has_paras=TRUE</code> ;
<code>tol</code>	A tolerance in absolute percent change in the global mean as the iterations progress. This tolerance should be good enough for many practical large sample applications, but can be reduced considerably if the function is used as the backdoor to integration of a given copula family instead of a sample estimator;
<code>minit</code>	Minimum number of iterations regardless of <code>tol</code> setting;
<code>maxit</code>	Maximum number of iterations;
<code>subdivisions</code>	Number of subdivisions and somewhat an analog to the term used in numerical integration. This subdivisions will be the size of the Monte Carlo sample for each iteration;
<code>large_n</code>	For an empirical copula and as the sample size gets large, not only does the σ_C computation slow down because a double summation is involved, but the counting that empirical copula must also perform gets large, therefore, the <code>para</code> that will be ultimately passed to the empirical copula is randomly subsetting by <code>large_n</code> . This is done in order to speed up iteration times at expense of increasing time that <code>tol</code> presumably is reached;
<code>include1moms</code>	A logical to include up to the first five linear moments for σ_C and logit-transformed $\log[\sigma_C/(1 - \sigma_C)]$ in the returned list.
<code>trace</code>	A logical, if true, that will produce a scatter plot of the sample (or the current iteration subset [refer to <code>large_n</code>]) along with the random sample points used for the current Monte Carlo iteration;
<code>verbose</code>	A logical providing percent change results by iteration to the console; and
<code>...</code>	Additional arguments to pass to the <code>halton()</code> function of the randtoolbox and the copula specified by <code>cop</code> (EMPIRCop) or a copula family.

Value

A list of the Monte Carlo integration results is returned containing

wolves	A vector of the σ_C by iteration;
estimates	The minimum, mean (the formal estimate $\hat{\sigma}_C$), maximum along with optional variance (computed from second L-moment), L-scale (second L-moment), L-skew, and L-kurtosis of the wolves;
estimates	Using logit-transformation of the wolves, the minimum, mean, maximum along with optional variance (computed from second L-moment), L-scale (second L-moment), L-skew, and L-kurtosis of the wolves;
its	Number of iterations before termination by maxits or upon reaching tol;
lastpctchg	The percent change on the last iteration.

Note

Beyond about 1,000 sample sizes `wolfCOP(as.sample=TRUE)` scales in time poorly, so we see in the following recipe the giant increase in speed at a small cost in precision on the Schweizer–Wolff Sigma estimate.

```
UV      <- simCOP(3000, cop=CLcop, para=3, graphics=FALSE, seed=473)
thewolf <- wolfCOP(      cop=CLcop, para=3) # 0.7864391
system.time( sfwolf <- wolfCOP(      para=UV, as.sample=TRUE,
                                     nlarge=1, usefastgrid=TRUE ) )
system.time( smwolf <- wolfCOP(      para=UV, as.sample=TRUE,
                                     nlarge=1, usefastgrid=FALSE) )
system.time( mcwolf <- wolfCOPsamc(para=UV,  usetime=TRUE) )
# user system elapsed   WOLF(sample fast) = 0.8002883
# 0.600  0.017  0.616
# user system elapsed   WOLF(sample slow) = 0.8002883
# 188.072 25.594 213.632
# user system elapsed   WOLF(monte carlo) = 0.799808
# 12.729  1.449 14.191
```

If one wanted to make some comparisons of computed results between `wolfCOP` and `wolfCOPsamc`, the following recipe, with a suitably large sample size, that we see a general spread around the equal value line. The use of Monte Carlo integration approach is not advised when computational speed of `wolfCOP` is sufficient. This `wolfCOPsamc` function is potentially of less usefulness than suggested because of the `EMPIRgrid_fast` within `wolfCOP` for rapid estimation by a gridded empirical copula.

```
nsam <- 1000; nsim <- 20; wolf <- mcwolf <- rep(NA, length=nsim)
for(i in seq_len(nsim)) {
  uv <- as.data.frame(matrix(runif(nsam*2), ncol=2))
  wolf[i] <- wolfCOP(      para=uv, as.sample=TRUE)
  mcwolf[i] <- wolfCOPsamc(para=uv)$estimates[2]
  print(c(i, wolf[i], mcwolf[i]))
}
df <- data.frame(sigma=wolf, sigma_mc=mcwolf)
df <- df[complete.cases(df),]
```

```

xylim <- range( c(range(df$sigma, na.rm=TRUE), range(df$sigma_mc)) )
plot(df$sigma, df$sigma_mc, xlim=xylim, ylim=xylim,
      xlab="Schweizer-Wolff sigma by whole-sample counting",
      ylab="Schweizer-Wolff sigma by Monte Carlo integration")
lines(par()$usr[1:2], par()$usr[3:4]) # equal value line

```

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

Schweizer, B., and Wolff, E.F., 1981, On nonparametric measures of dependence for random variables: The Annals of Statistics, v. 9, no. 4, pp. 879–855, <https://www.jstor.org/stable/2240856>.

See Also

[wolfCOP](#), [EMPIRCop](#)

Examples

```

n <- 40; para=c(1.3, 2.4)           # Small sample, for speed.
UV <- simCOP(40, para=para, cop=GHCop, seed=1) # Simulate a sample
wolfCOP( para=para, cop=GHCop)      # Theo. value + next 2 lines
wolfCOPsamc(para=para, cop=GHCop)$estimate[2] # Same as next line
wolfCOPsamc(para=para, cop=GHCop, para_has_paras=TRUE)$estimate[2]
wolfCOP( para=UV, as.sample=TRUE, usefastgrid=TRUE, nlarge=1) # Spring 2026
wolfCOP( para=UV, as.sample=TRUE, usefastgrid=FALSE) # original pkg implementation
wolfCOPsamc(para=UV)$estimate[2]    # Monte Carlo estimate

## Not run:
# Test the trace and verbose arguments
para <- list(cop=c(P, M), part=c(0,0.5,1))
UV <- simCOP(1000, cop=ORDSUWcop, para=para)
SW <- wolfCOPsamc(para=UV, tol=0.0005, trace=TRUE, verbose=TRUE) #
## End(Not run)

## Not run:
# Test the various interfaces for a sample computation along with
# setting the usetime to the random number subsystem.
UV <- simCOP(1000, cop=CLcop, para=3, graphics=FALSE)
mc1 <- wolfCOPsamc(para=UV, usetime=TRUE)$estimates[2]
mc2 <- wolfCOPsamc(u=UV, usetime=TRUE)$estimates[2]
mc3 <- wolfCOPsamc(u=UV[,1], v=UV[,2], usetime=TRUE)$estimates[2] #
## End(Not run)

## Not run:
# Let us check on the veracity of wolfCOPsamc() by a cluster-based computation
# of realization of mean values of Lmoments of logit-transformed Sigmas for a

```

```

# given sample size against the projected Lmoments (Lskew and Lkurtosis) in
# regressions implemented within wolfCOPtest(). We will need the ellipse and
# parallel packages already installed to the user's machine for this test.
nsam <- 40 # Sample size (not related to simulation settings in the example.)
lmomco::plotlrmrdia(lmomco::lrmrdia(), empty=TRUE, # Lmoment ratio diagram
  xlim=c(0.1, 0.35), ylim=c(0.1, 0.20), las=1, xaxs="i", yaxs="i")
lmomco::plotlrmrdia(lmomco::lrmrdia(usrtrim=TRUE), add=TRUE, expand.names=TRUE,
  autolegend=TRUE, xleg="topleft", ncol=1, nolimits=TRUE,
  nogov=TRUE, nogpa=TRUE, noexp=TRUE, nonor=TRUE, nogum=TRUE,
  noray=TRUE, nouni=TRUE, noaep4=TRUE, noglo=TRUE, nopdq3=TRUE)
xy <- NULL # coordinates of the Lskew (T3) and Lkurtosis (T4) of logit-Sigma
for(n in c(3:99, 10^seq(2, 6, by=0.1))) { # a vector of sample sizes
  wt <- wolfCOPtest(0, n)$lmoms_logit_sigma # Lmoments of logit-Sigma
  xy <- rbind(xy, data.frame(tau3=wt[3], tau4=wt[4])) # isolate the T3 and T4
}
lines( xy[,1], xy[,2], col="wheat3", lwd=4) # plot the estimation of T3 and T4
wt <- wolfCOPtest(0, nsam)$lmoms_logit_sigma # isolate the prediction for the
points(wt[3], wt[4], pch=16) # sample size that we are studying
cols <- c("salmon4", "salmon1", "seagreen4", "seagreen1")
# All the code below can be run with on setting of doMC false and then true.
doMC <- FALSE # set to true for wolfCOPsamc() else wolfCOP(as.sample=TRUE)
LMR <- NULL; ncore <- 8; nloop <- 40; nsim <- 500 # last of initializations
# The wolfCOPsamc() has long running time but by about 500-1000 samples it
# likely will then begin outpacing the wolfCOP(). We have the nsam default as
# small for this example and nsim enough for definitive stability in the results
# and nloop large enough for an authoritative ellipse to be drawn.
if(nloop < 4) stop("Need at least four for Lmoment computation later")
for(j in seq_len(nloop)) {
  message(j, "-", appendLF=FALSE)
  if(length(grep("[02468]0$", as.character(j))) > 0) message("", appendLF=TRUE)
  cl <- parallel::makeCluster(getOption("cl.cores", ncore)) # make CPU cluster
  parallel::clusterExport( cl, "nsam") # make "nsam" visible in cluster FUN
  if(doMC) { # set tolerance "up" just for some speed increase for the example
    wolves <- parallel::parSapply(cl=cl, seq_len(nsim), FUN=function(i) {
      wolfCOPsamc(para=as.data.frame(matrix(runif(nsam*2),
        ncol=2)), tol=0.01, usetime=TRUE)$estimates[2] })
  } else {
    wolves <- parallel::parSapply(cl=cl, seq_len(nsim), FUN=function(i) {
      wolfCOP( para=as.data.frame(matrix(runif(nsam*2),
        ncol=2)), as.sample=TRUE)
    })
  }
  parallel::stopCluster(cl) # stop the cluster as it will be restarted as needed
  wolves <- log( wolves / (1 - wolves) ) # make logit-Sigmas
  lmr <- Lmoments::Lmoments(wolves) # faster than lmomco::lmoms()
  lmr[3] <- lmr[3] / lmr[2]; lmr[4] <- lmr[4] / lmr[2] # Lskew and Lkurtosis
  LMR <- rbind(LMR, data.frame(L1=lmr[1], L2=lmr[2], T3=lmr[3], T4=lmr[4]))
  if((par()$usr[1] <= lmr[3] & lmr[3] <= par()$usr[2]) &
    (par()$usr[3] <= lmr[4] & lmr[4] <= par()$usr[4])) {
    if(doMC) {
      points(lmr[3], lmr[4], col=cols[1], bg=cols[2], pch=24, lwd=0.9, cex=0.8)
    } else {
      points(lmr[3], lmr[4], col=cols[3], bg=cols[4], pch=25, lwd=0.9, cex=0.8)
    }
  }
}

```

```

    }
  }
  # Lskew and Lkurtosis are approximately multivariate normally distributed.
  Trho <- stats::cor( LMR$T3, LMR$T4 ) # Pearson correlation coefficient
  Tsc1 <- c(Lmoments::Lmoments(LMR$T3)[2] * sqrt(pi), # standard deviations in U
           Lmoments::Lmoments(LMR$T4)[2] * sqrt(pi)) # standard deviations in V
  Tctr <- c(mean(LMR$T3), mean(LMR$T4)) # ellipse center
  Telp <- ellipse::ellipse(Trho, scale=Tsc1, centre=Tctr, npoints=200, level=0.95)
  if(doMC) {
    lines(Telp, col=cols[1], lty=4, lwd=2) # ellipse itself
    points(mean(LMR$T3), mean(LMR$T4), pch=21, cex=2, col=cols[3], bg=cols[2])
  } else {
    lines(Telp, col=cols[3], lty=4, lwd=2) # ellipse itself
    points(mean(LMR$T3), mean(LMR$T4), pch=21, cex=2, col=cols[3], bg=cols[4])
  }
  points(wt[3], wt[4], pch=16) # sample size that we are studying #
## End(Not run)

```

wolfCOPtest

Empirically Created Schweizer and Wolff Sigma Test Against an Independence Copula

Description

Perform an *Independence test* based on a *Schweizer–Wolff Sigma* test for σ_C stemming from a Null hypothesis of $\text{NULL} : \mathbf{C} = \mathbf{\Pi}$ (*Independence copula*, [P](#)), where σ_C is defined and computed under [wolfCOP](#). There are no other known hypothesis tests tied to Schweizer–Wolff Sigma σ_C in the literature.

This *Schweizer–Wolff Sigma hypothesis test* in **copBasic** has been empirically tuned from small to large samples by extensive simulation, and, as implemented, the test is closely tied to the version of sample computation by [wolfCOP](#) using a particular form of the empirical copula (refer to that function for details). The `wolfCOPtest` function internally loads the `wolfCOPtest_data_smlsam` tables, and most users generally will not need to access that those tables themselves unless perhaps for checking veracity of the simulations.

Usage

```

wolfCOPtest(x, y, asuv=FALSE, aslist=TRUE, na.rm=TRUE, digits=6,
            probs=c(0.90, 0.95, 0.98, 0.99, 0.995),
            zmat=NULL, statf=mean, rndphi=20, usepade=FALSE,
            ties.method=c("average", "first", "last", "random", "max", "min"), ...)

```

Arguments

x Nonexceedance probabilities u in the X direction if `asuv=TRUE`, else data in the X direction for which nonexceedance probabilities will be computed by plotting-position formula. If `x` is a single value, it is assumed to be a previously computed σ_C and `y` will be considered as the sample size. Note, statistical

	estimation is based on samples of $n \geq 3$, if the sample size is less than this, then internally $n = 3$ is set for the computations. The x can also be a two column data structure and the y will be taken from the second column;
<code>y</code>	Nonexceedance probabilities v in the Y direction if <code>asuv=TRUE</code> , else data in the Y direction for which nonexceedance probabilities will be computed by plotting-position formula (refer to <code>lmomco:pp()</code> function, but simply the <code>rank(...)</code> function is used);
<code>asuv</code>	Logical identifying whether the x and y are already in nonexceedance probabilities or rather such need to be computed internally by plotting-position formula (refer to <code>lmomco:pp()</code> function, but simply the <code>rank(...)</code> function is used);
<code>aslist</code>	Logical to return a list of the test results rather than a vector;
<code>na.rm</code>	A logical to trigger a <code>complete.cases</code> operation on internal matrix of the data as probabilities;
<code>digits</code>	Digits for rounding before return;
<code>probs</code>	Quantiles by nonexceedance probability to extract from the fitted distribution—these are common upper-tail critical values. If <code>probs</code> is empty, then 0.95 nonexceedance probability is silently used;
<code>zmat</code>	An optional matrix of four columns and rows equally those of the data values with columns 1 and 2 containing, respectively, the lower and upper limits of the u and columns 3 and 4 containing, respectively, the lower and upper limits of v . This <code>zmat</code> feature provides for support if data censoring is involved and simulation between the limits made (refer to Examples). Note, if <code>zmat</code> is given, then <code>ties.method</code> is set "random" and <code>asuv</code> is set false;
<code>statf</code>	Statistical function for reducing simulated σ values (mean or median suggested), but a user could <code>plot(density(result\$rand_sigma))</code> and reach their own decisions on centrality measure;
<code>rndphi</code>	A multiplier on the number of ties or censored values to drive simulation of samples as part of populating the <code>rand_sigma</code> element, when applicable, on the function return;
<code>usepade</code>	A logical to switch, on estimation of the empirical L-moments of the logit transformed simulations, between (1) power-exponent optimized regression equations, or (2) optimized <i>Padé Approximant</i> forms;
<code>ties.method</code>	Argument of the same name for <code>rank</code> , but if there are ties, <code>asuv</code> is set false, and <code>ties.method</code> is set to random, then simulation is made internally on the ranking and <code>statf</code> used to compute the σ estimate;
<code>...</code>	Additional arguments to pass to the <code>lmomco:pp()</code> function.

Details

The implementation here in **copBasic** to approximate p-values is based on *Monte Carlo simulations* of Π and computation of the L-moments of a *logit transform* of the computed σ_Π for which non-linear regression of the L-moments to the sample size have been created. Development scripts for the simulations are located in the subdirectory `inst/make_wolfCOPtest/` of **copBasic** sources.

The *logit transformation* is defined as

$$\text{logit}(\sigma_{\mathbf{C}}) = \log\left(\frac{\sigma_{\mathbf{C}}}{1 - \sigma_{\mathbf{C}}}\right) = \eta,$$

and its inverse is defined as

$$\sigma_{\mathbf{C}} = \text{logit}^{(-1)}(\eta) = \frac{1}{1 + \exp(-\eta)}.$$

Sample-sample empirical distributions ($n \in 3, \dots, 40$) will have p-values reported on output and missing for $n \geq 41$ (refer to [wolfCOPtest_data_sm1sam](#) for more details).

The predicted L-moments are used to drive from the **lmomco** package the Pearson type III (parpe3, cdfpe3, quape3) of the *logit transform* of $\sigma_{\mathbf{\Pi}}$ to compute the p-value and supporting information.

Evaluation of an *L-moment ratio diagram* (refer to **lmomco** functions `plotlmrda()` and `lmrda()`) shows a change from a Generalized Normal distribution to Pearson Type III at about $n = 40$, but inspection of upper-tail empirical distributions (and hence p-values in the practical range for NULL rejection) visually suggest that the Pearson Type III upper tail is better choice for overall application. The `wolfCOPtest` function, therefore, exclusively uses the Pearson Type III until further evidence were to manifest.

To conclude, the figure below shows the *L-moment ratio diagram* of the *L-skew* and *L-kurtosis* diagram for the logit transform of the *L-moments* of the $\hat{\sigma}_{\mathbf{C}}$ based on [EMPIRgrid_fast](#) in the aforementioned build directory of the **copBasic** package. The figure is based on nearly 900 million simulations summed across a representative range of sample sizes $n \in 8, \dots, 15001$, and provides support for the decision by the author to use the Pearson Type III as the distribution model for the hypothesis test.

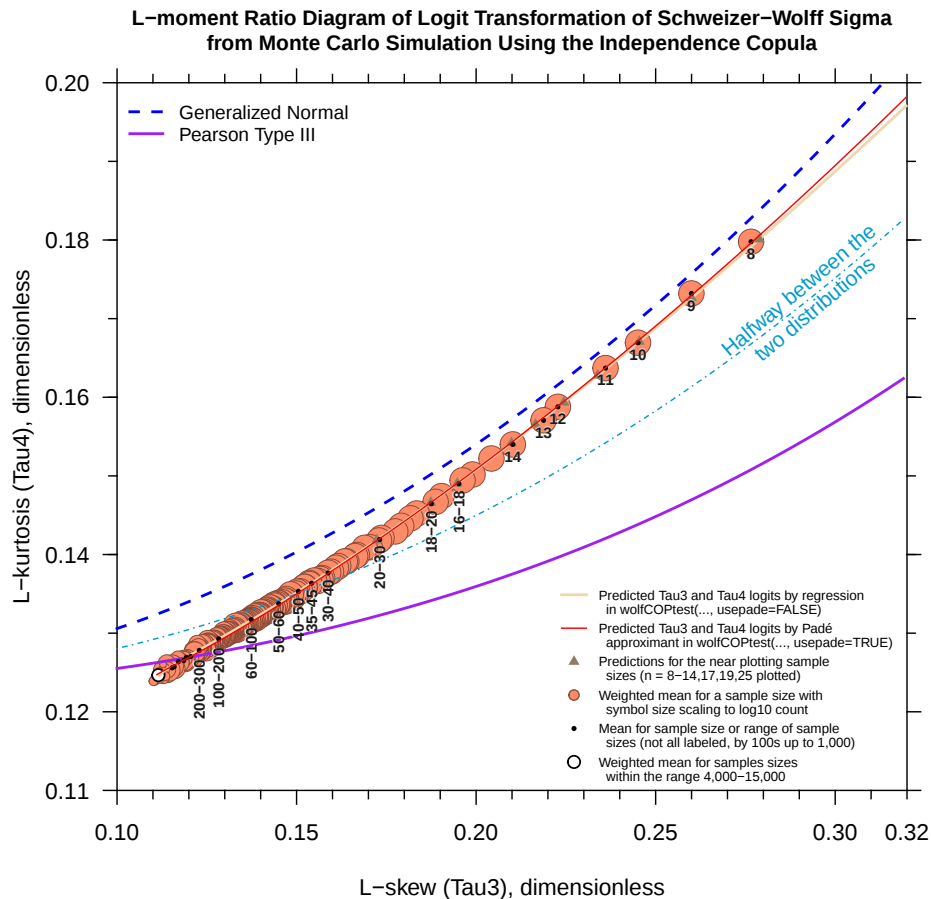


Figure. L-moment ratio diagram of simulated values for *Schweizer–Wolff Sigma* showing trajectory transitioning from near a Generalized Normal to a Pearson Type III as sample size increases along with parametric equations fits implemented for the Hypothesis Test

Value

Extensive list or named vector output the test is returned. For the list, the estimate contains duplicate information as `statistic[1]`; this is so that there is an estimate entry to parallel the return of `cor.test(...)` to help users write nomenclature-similar calls between the functions.

Author(s)

W.H. Asquith

See Also

[wolfCOP](#), [wolfCOPsamc](#), [wolfCOPtest_data_smlsam](#)

Examples

```

uv <- simCOP(n=62, cop=PSP, graphics=FALSE, seed=8)
# Approx p-value 0.001 PSP is not independence.
# There is no p.value(sample_le40) entry on the return because too large sample size.
wolfCOPtest(uv[,1], uv[,2], asuv=TRUE, aslist=FALSE, usepade=FALSE)
wolfCOPtest(uv[,1], uv[,2], asuv=TRUE, aslist=FALSE, usepade=TRUE )
# See how the p-values between the empirical approximations are about the same.

uv <- simCOP(n=62, cop=P, graphics=FALSE, seed=8)
wolfCOPtest(uv[,1], uv[,2], asuv=TRUE, aslist=FALSE) # approx p-value 0.200***
# Note, the small sample p-values are NA for two previous examples because the sample
# size has become "large", but in the next example, we can inspect two p-values
# (1) by the Pearson type III distribution to the predicted L-moments for the sample
# size and (2) by the small-sample empirical tables in object wolfCOPtest_data_smlsam.
uv <- simCOP(n=13, cop=P, graphics=FALSE, seed=81)
wolfCOPtest(uv[,1], uv[,2], asuv=TRUE, aslist=FALSE)
# p.value(dist_pe3) p.value(sample_le40) # four signif. figures kept here as
# 0.2550*** 0.25144** # simulations might tune up internals

## Not run:
# This example needs larger nsim but will show that the Sigma is performing close to
# Spearman Rho, which helps establish the veracity of the algorithms. If the user were
# to switch out the copula P and replace with CIRCcop (circular copula), then one will
# see Sigma performing similarly but Rho failing because the CIRCcop is an example of
# perfect dependence in a setting of independence. Use of much larger nsim for this
# example will hone the apparent rejection to about 0.05 for P, but clearly we see
# divergence in performance for CIRCcop.
alpha <- 0.05; n <- 42; nsim <- 20000; cop <- P; p <- NULL
set.seed(1)
wolf <- replicate(nsim, { wolfCOP(para=rCOP(n=n, cop, para=p),
                                as.sample=TRUE, nlarge=0, usefastgrid=TRUE) })
wolfpv <- sapply(wolf, function(w) {
  wolfCOPtest(w, n, asuv=TRUE, aslist=FALSE)[4]})
message(" Schweizer-Wolff Sigma(n=", n, ") NULL=P rejection rate = ",
        sum(wolfpv < alpha) / nsim)
set.seed(1)
rhopv <- replicate(nsim, { uv <- rCOP(n=n, cop=cop, para=p);
  cor.test(uv[,1], uv[,2], method="spearman")$p.value })
message(" Spearman Rho(n=", n, ") NULL=P rejection rate = ",
        sum(rhopv < alpha) / nsim)
# For nsim <- 20000
# P : Schweizer-Wolff Sigma(n=42) NULL=P rejection rate = 0.04935
# P : Spearman Rho(n=42) NULL=P rejection rate = 0.04805
# CIRCcop : Schweizer-Wolff Sigma(n=42) NULL=P rejection rate = 0.28125
# CIRCcop : Spearman Rho(n=42) NULL=P rejection rate = 0.00100
# Such example computations show that Rho does not "sense" the perfect dependence
# of CIRCcop because its "correlation" is zero but has obvious dependence when
# plotted. Therefore, Sigma has a much higher rejection rate that will go to unity
# as the sample size increase.
## End(Not run)

## Not run:

```

```

lmomco::plotlmr dia(lmomco::lmr dia(usrtrim=TRUE), nopoints=TRUE, xleg="topleft",
                    xlim=c(0.11,0.32), ylim=c(0.12,0.20), autolegend=TRUE,
                    expand.names=TRUE, autoaxes=TRUE, xaxs="i", yaxs="i", lwd.cex=1.3)
xy <- st <- NULL
ns <- sort(unique(c(10^seq(0,1,0.01), 10^seq(0, 4, by=0.02))))
for(n in ns) {
  lmr <- wolfCOPtest(0, n, usepade=FALSE)$lmoms_logit_sigma; if(is.null(lmr)) next
  xy <- rbind(xy, data.frame(n=n, mu=lmr[1], lam2=lmr[2], tau3=lmr[3], tau4=lmr[4]))
}
xy <- xy[par()$usr[1] <= xy$tau3 & xy$tau3 <= par()$usr[2],]
xy <- xy[par()$usr[3] <= xy$tau4 & xy$tau4 <= par()$usr[4],]
lines(xy$tau3, xy$tau4, col="grey60", lwd=7)
for(n in ns) {
  lmr <- wolfCOPtest(0, n, usepade=TRUE)$lmoms_logit_sigma; if(is.null(lmr)) next
  st <- rbind(st, data.frame(n=n, mu=lmr[1], lam2=lmr[2], tau3=lmr[3], tau4=lmr[4]))
}
st <- st[par()$usr[1] <= st$tau3 & st$tau3 <= par()$usr[2],]
st <- st[par()$usr[3] <= st$tau4 & st$tau4 <= par()$usr[4],]
lines(st$tau3, st$tau4, col="red1", lwd=2)
txt <- c("Logit transform sampling Sigma distribution, wolfCOPtest(usepade=FALSE)",
        "Logit transform sampling Sigma distribution, wolfCOPtest(usepade=TRUE)")
legend("bottomright", txt, lwd=c(7,2), col=c("grey50", "red1"), bty="n", cex=0.9) #
## End(Not run)

## Not run:
# Here is a cross check in properties of the mathematics computing W.
# The examples here also show the pass of 2-columns in x; so, a local copy
# of a simulations is not needed before passing to wolfCOPtest().
alpha <- 0.05; n <- 42; nsim <- 2000; cop <- P; p <- NULL
set.seed(1)
wolf1 <- replicate(nsim, { wolfCOP(para=rCOP(n=n, cop, para=p),
                                   as.sample=TRUE, nlarge=0, usefastgrid=TRUE) })
wolf1pv <- sapply(wolf1, function(w) { # here, sigma precomputed
  wolfCOPtest(w, n, asuv=TRUE, aslist=FALSE)[4]})
set.seed(1)
wolf2pv <- replicate(nsim, { # here the "true" probabilities are known
  wolfCOPtest(rCOP(n=n, cop, para=p), asuv=TRUE)$p.value[1]})
set.seed(1)
wolf3pv <- replicate(nsim, { # here the probabilities computed internally
  wolfCOPtest(rCOP(n=n, cop, para=p), asuv=FALSE)$p.value[1]})
message(" Schweizer-Wolff Sigma(n=", n, ") NULL: P rejection rate = ",
        sum(wolf1pv < alpha) / nsim)
message(" Schweizer-Wolff Sigma(n=", n, ") NULL: P rejection rate = ",
        sum(wolf2pv < alpha) / nsim)
message(" Schweizer-Wolff Sigma(n=", n, ") NULL: P rejection rate = ",
        sum(wolf3pv < alpha) / nsim) #
## End(Not run)

## Not run:
# A micro cross check between wolfCOPtest and Independence test of copula package
n <- 60; J <- copula::indepTestSim(n, p=2, N=100000, verbose=FALSE)
for(i in 1:20) {
  uv <- simCOP(n, cop=M0cop, para=c(0.2, 0.8), seed=i)

```

```

    pw <- wolfCOPtest(uv, asuv=FALSE)$p.value[1]
    pi <- copula::indepTest(uv, J)$pvalues[1]
    mtext(paste0("seed=", i, "; wolfCOPtest=", pw, "; indepTest=", pi))
  } #
## End(Not run)

## Not run:
# Here is a cross check of sorts between wolfCOPtest and the Independence test
# available in the copula package. We use a Clayton copula with small bivariate
# association and will see for this circumstance that wolfCOPtest out performs
# copula::indepTest by just a small amount by rejecting the NULL more often.
sample_size <- 100; ALPHA <- 0.05
THECOP <- CLcop; THEPARA <- 0.20; rhoCOP(cop=THECOP, para=0.20) # 0.1358466

nsim <- 2000; digits <- as.integer( floor( log10(nsim) ) )
J <- copula::indepTestSim(sample_size, 2, N=100000)

W <- I <- NULL
for(i in seq_len(nsim)) {
  if(is.null(W)) {
    uv <- copBasic::simCOP( sample_size, cop=THECOP, para=THEPARA )
    mtext("Copula simulation reference figure", line=0.5, font=2 )
  } else {
    uv <- copBasic::rCOP( sample_size, cop=THECOP, para=THEPARA )
  }
  I <- c(I, copula::indepTest( uv, J )$pvalues[1])
  W <- c(W, copBasic::wolfCOPtest(uv[,1], uv[,2], asuV=TRUE )$p.value[1])
}

par(lend=2, las=1); hyphen <- "\u00ad"
plot( sort( rank(I) ) / ( length(I) + 1 ), sort(I), type="l", xaxs="i", yaxs="i",
      xlab="Nonexceedance probability", lwd=3, col="grey50",
      ylab=paste0("P", hyphen, "value of respective test of independence"))
lines(sort( rank(W) ) / ( length(W) + 1 ), sort(W), col="darkorange1")
txt <- c(paste0("Independence test (copula::indepTest) has rejection rate = ",
               round( sum(I < ALPHA) / nsim, digits=digits), "%"),
        paste0("Schweizer-Wolff test (copBasic::wolfCOPtest) has rejection rate = ",
               round( sum(W < ALPHA) / nsim, digits=digits), "%"))
legend("topleft", txt, bty="o", box.lty=0, bg=NA, inset=0.02, cex=0.9,
      lwd=c(3,1), col=c("grey50", "darkorange1"))
mtext(paste0("Empirical distribution of p", hyphen, "values"), line=0.5, font=2) #
## End(Not run)

## Not run:
# DATA CENSORING EXAMPLE -----
set.seed(10) # Example using the zmat feature for support of censoring by simulation.
muslp <- 0.005; sds1p <- 0.019 # drift with exp used to keep parameter positive
# (1) Try rerunning the example moving sds1p up and down and review the results.
# (2) Try disabling the censoring for the 35+th observations and repeat (1).
yy <- 1980:2021; n <- length(yy) # Simulated log-Pearson III annual peak streamflows
qq <- 10^sapply(yy, function(y) { # put a small drift in the log10 mean and variation
  para <- lmomco::vec2par(c(3.5+(y-min(yy))*muslp,
                           exp(log(0.5)-(y-min(yy))*sds1p), 0.3), type="pe3")

```

```

      lmomco::rlmomco(1, para) }) # flow
zul <- zur <- rep(NA, n) # initialize the lower and upper limits for years
zvl <- zvr <- rep(NA, n) # initialize the lower and upper limits for flows
# NA years are not simulated and the data are used as is
# Make some highly contrived censoring for the 3, 6, 9, 13, and 35+th observations.
zul[ 3] <- yy[ 3] - 1.1; zur[ 3] <- yy[ 3] + 1.1 # Unusual to not know the year, but
zul[13] <- yy[13] - 1.5; zur[13] <- yy[13] + 2.4 # showing generalization as needed.
zvl[ 3] <- 10; zvr[3] <- qq[3]
zvl[ 6] <- qq[6]; zvr[6] <- max(qq) + sqrt(.Machine$double.eps) # nudge above max
zvl[ 9] <- 10^(log10(qq[9]) - 0.3); zvr[9] <- 10^(log10(qq[9]) + 0.2) # log-cycles
qq[ 9] <- sqrt( zvl[9] * zvr[9] ) # geometric mean
zvl[(n-6):n] <- 10; zvr[(n-6):n] <- qq[(n-6):n]

xy <- cbind(yy, qq); zmat <- cbind(zul, zur, zvl, zvr)
swolf <- wolfCOPtest(xy[,1], xy[,2], zmat=zmat, asuv=FALSE, statf=median, rndphi=500)
ylim <- log10( range( qq ) ); ylim <- 10^c( floor( ylim[1] ), ceiling( ylim[2] ) )
plot( yy, qq, type="n", log="y", xlab="Water year", ylab="Streamflow", ylim=ylim)
segments(x=zul, x1=zur, y=qq, y1=qq, col="darkgoldenrod3")
segments(x=yy, x1=yy, y=zvl, y1=zvr, col="darkgoldenrod3")
points(yy, qq, bg="darkgoldenrod1", col="darkgoldenrod4", pch=21)
naive <- wolfCOPtest(xy[,1], xy[,2], asuv=FALSE, ties.method="random" )
txt <- paste0("Sigma simulation (nsim=",
              prettyNum(length(swolf$rand_sigma), big.mark=","), ") ",
              "p.value=", round(swolf$p.value[1], digits=3), "; ",
              "Naive p.value=", round(naive$p.value[1], digits=3))
mtext(txt, line=1) # Visually the point values tell us to expect independence to be
# rejected, but when the data censoring is taken into account and simulation used,
# we do not reject independence that is heavily influenced by the last seven data
# values on the right being censored down through the bottom axis.
swolf$rand_sigma <- sort(swolf$rand_sigma)
p <- sapply(swolf$rand_sigma, function(k) { # distribution of p-values
  n <- ifelse(swolf$sample_size <= 40, 2, 1)
  wolfCOPtest(k, swolf$sample_size)$p.value[n] })
names(p) <- NULL; swolf$rand_p.values <- p
# plot(swolf$rand_sigma, swolf$rand_p.values, pch=16, col=grey(0, 0.05)) #
## End(Not run)

```

wolfCOPtest_check

*Empirical Comparison by Simulation between Independence Test by
Schweizer–Wolff Sigma from this package to Independence Test from
copula Package*

Description

Report on a simulation study of empirical performance an *Independence test* based on a *Schweizer–Wolff Sigma* test for σ_C stemming from a Null hypothesis of $\text{NULL} : \mathbf{C} = \mathbf{\Pi} = uv$ (*Independence copula*, [P](#)), where σ_C is defined and computed under [wolfCOP](#). The study uses a *Clayton copula* ([CLcop](#)) is $C_{\Theta}(u, v)$ with parameter Θ set through *Kendall Tau* ([tauCOP](#)) by $\tau = \Theta/(\Theta + 2)$ and $\Theta = 2\tau/(1 - \tau)$.

The **copBasic** package provides an *Independence test* with approximated p-values in **wolfCOPtest**, and the **copula** package provides an *Independence test* by **copula** function `indepTest`. Both tests can detect dependency even if *Kendall Tau* and *Spearman Rho* are equal to zero. Example of comparison between the two tests is in the **Examples** in **wolfCOPtest**.

The **wolfCOP** uses *Schweizer–Wolff Sigma* σ_C of a copula according to Nelsen (2006, p. 209) by

$$\sigma_C = 12 \int \int_{T^2} |\mathbf{C}(u, v) - uv| \, du dv,$$

which is $0 \leq \sigma_C \leq 1$. The **wolfCOPtest** is based on simulation of a sampling estimator of σ_C and detailed parameterization of a three-parameter probability distribution by the *method of L-moments*. The sampling estimator used (refer to **wolfCOP**) was

$$\hat{\sigma}_C = \frac{12}{n^2 - 1} \sum_{i=1}^n \sum_{j=1}^n \left| \hat{\mathbf{C}}_n\left(\frac{i}{n}, \frac{j}{n}\right) - \frac{i}{n} \times \frac{j}{n} \right|,$$

where $\hat{\mathbf{C}}_n$ is the simplest empirical copula of

$$\hat{\mathbf{C}}_n\left(\frac{i}{n}, \frac{j}{n}\right) = \frac{1}{n} [\# \text{ of } (U_k \leq U_i, V_k \leq V_j)].$$

The **copula** package in function `indepTestSim` is based around the test statistic (as a bivariate form) (Hofert *et al.*, 2017, p. 174) (for such a form refer also to **hoeFCOP**)

$$S_n^\Pi = \int \int_{T^2} n(\mathbf{C}_n(u, v) - uv)^2 \, du dv,$$

where $\mathbf{C}_n(u, v)$ is an empirical copula.

Originating literature leading to S_n^Π through citation by Hofert *et al.*, (2017, p. 173–178) are Deheuvels (1981), Genest and Rémillard (2004), and Genest *et al.* (2007). Genest and Rémillard (2004) show, repeated in Hofert *et al.* (2017, p. 175), and implemented in **copula** function `indepTest` that approximate p-values can be computed by simulation of size N by

$$\frac{1}{N+1} \left(\sum_{k=1}^N \mathbf{1}(S_n^{\Pi, (k)} \geq S_n^\Pi) + \frac{1}{2} \right),$$

for S_n^Π being a large simulation of the Π copula. The simulation study described in the next section, used a sample size of $N = 100,000$ for the **copula** package function `J <- indepTestSim(n, 2, N=100000)` where n is the sample size of the pending test data. This `J` object was used for the `indepTest` calls. This is a required step unique to **copula** package implementation as described by Hofert *et al.* (2017, p. 175) and package documentation. So, the N simulation size in the aforementioned equation is the simulation size on which `indepTest` bases its p-values and therefore is unique to needs of the **copula** package.

This Simulation Study—A sequence of $\tau \in (0, 0.02, \dots, 0.48, 0.50)$ were converted to parameters Θ of the *Clayton copula* (**CLcop**). Values for τ were selected to start the $\mathbf{C}_{\Theta=0} \equiv \Pi$, which provides a chance to study results with the condition of the Null being Π . The purpose is to compare for a range of sample sizes (n) the rejection performance of **wolfCOPtest** to the **copula** function

indepTest. The script is provided in `inst/make_wolfCOPtest/chck_wolfCOPtestCL.R` of this **copBasic** package source code.

For each of the τ values and hence *Clayton copula* parameter Θ values, a Π sample is drawn for each of sample sequences of sizes $n \in (5, 20; 1), (22, 50; 2), (55, 100; 5)$. For a given sample, the **wolfCOPtest** and indepTest were made and p-values compared to the $\alpha = 0.05$ statistical significance level; a binary then is recorded as whether the null hypothesis of independence is rejected or not.

The aforementioned process was repeated for large simulation sizes of $m = 150,000$ for all τ used (total simulation $\sum m = 6,519,000$). The differences in simulation size are heuristic to enhance the precision of the results for small τ values. In the end, for each of the τ for each of the n for simulation size m , the total number of rejections of the null hypothesis was counted for each method and divided by the number of simulations, which converts the binary into a rejection rate. The rejection rate between methods at the $\alpha = 0.05$ significance level becomes the measure on which to assess performance of **wolfCOPtest** to indepTest.

The relation between simulated rejection rates for independence at $\alpha = 0.05$ significance level for a selected subset of $\tau \in (0.00, 0.48; 0.04)$ is shown in figure 1. The τ were subsetting for graphical reasons. The lower left corner as a zoom region demarcated that is shown in figure 2. Each of the two figures as the $\alpha = 0.05$ significance level drawn as a horizontal line.

The figures have the rejection rates for **wolfCOPtest** drawn as solid lines and rejection rates for indepTest drawn as the dashed lines. The line color ramps with the τ value and these Kendall Taus are labeled and value plotted between the two curves. The two curve types show that both methods have a similarity of performance. On scrutiny, the curves for **wolfCOPtest** generally plot just above those for the indepTest. This is an indication of higher rejection rates of the null hypothesis as departure from independence ($\tau > 0$) seems to increase but only for a range of sample sizes. Figure 1 shows that as sample sizes become large relative to the τ is a convergence towards 100-percent rejection (a rejection rate = 1).

Figure 2 is a zoom into the lower left corner of figure 1 and is an emphasis of the method performance for very small sample sizes. It appears that indepTest for $\tau \leq 0.02$ barely outperforms **wolfCOPtest**. A general conclusion, however, from the figures is that the **wolfCOPtest** by **copBasic** is either better tuned empirically than how indepTest is implemented by **copula** or that the absolute value operation (difference between empirical copula and $\Pi = uv$) in the definition of σ_C in lieu of the square operation (squared difference between empirical copula and $\Pi = uv$) in S_n^Π .

The mean difference between **wolfCOPtest** and indepTest rejection is about 1 percent for the $\tau \in (0, 0.02, \dots, 0.48, 0.50)$ with $n \in (5, 20; 1), (22, 50; 2), (55, 100; 5)$ totalling about 3.7 million simulations. This is evidence that both methods perform similar to each other, which provides for documentation such as this for **copBasic** that the *Schweizer–Wolff Sigma* test is well implemented beyond the evidence documented in **Examples** of **wolfCOPtest**.

The results in figures 1 and 2 are compelling. Both figures as exemplified by empirical simulation that **wolfCOPtest** does outperform the independence test of indepTest as implemented within the **copula** package. Though the rejection rates of *Kendall Tau* or *Spearman Rho* are not otherwise indicted in those two figures (figs. 1 and 2), the practitioner could study the scripted results of `inst/make_wolfCOPtest/chck_wolfCOPtestCIRC.R`, which is built around application of the *Circula copula* (**CIRCcop**) to further explore the performance of the *Schweizer–Wolff Sigma* test. Those results for $m = 150,000$ are shown in figure 3 and show for that copula (**CIRCcop**) that **wolfCOPtest** continues to outperform indepTest in terms of rejection at the 0.05 significance level.

Figure 3 also has highlighted at $50\times$ the computed rejection of independence, the trajectory of the rejection of independence from the Circular copula by well-known statistics of *Kendall tau* and *Spearman Rho*. The trajectories of the simulations for those two statistics require a multiplication of 50 to avoid changes in vertical scale and plotting of a second vertical axis on the right.

A quite curious aspect is that the rejection rates dip below the $\alpha = 0.05$ for sample sizes $n \in 6, \dots, 19$ and possibly this is related to algorithmic estimation of nonexceedance probabilities for small samples. Beyond $n = 20$ both methods reject with increasing regularity of independence. By $n \approx 80$ but methods are rejecting independence at 100 percent. However, within the sample range of about $n \in (20+, \dots, 80)$, **wolfCOPtest** is superior.

Ultimately, it is not known for all settings or sample size whether or not **wolfCOPtest** will show to be generally more precise at rejection of *Independence* for all sample sizes. This assertion is based entirely on the heuristics of empirical simulation. Evidence is provided, however, in this **copBasic** documentation that the implementation of a *Schweizer–Wolff Sigma* test appears to have marginally better performance than **copula** package `indepTest`.

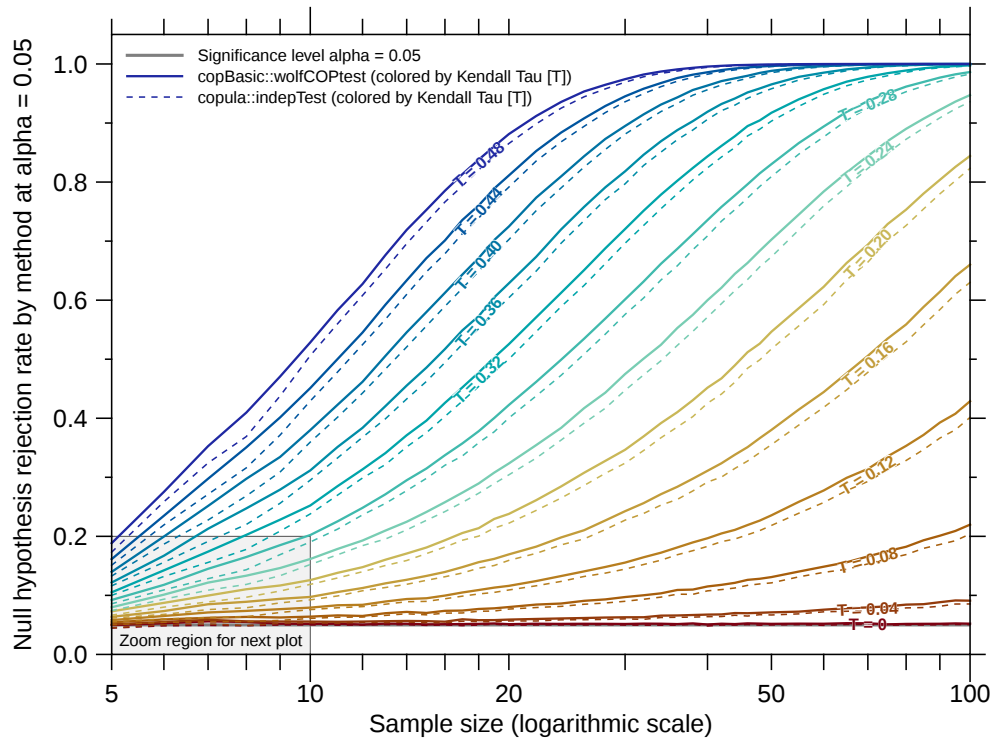


Figure 1. Relation between simulated rejection rates for independence at $\alpha = 0.05$ significance level for selected sample sizes 5–100 for a Clayton copula parameterized by given Kendall Tau as indicated

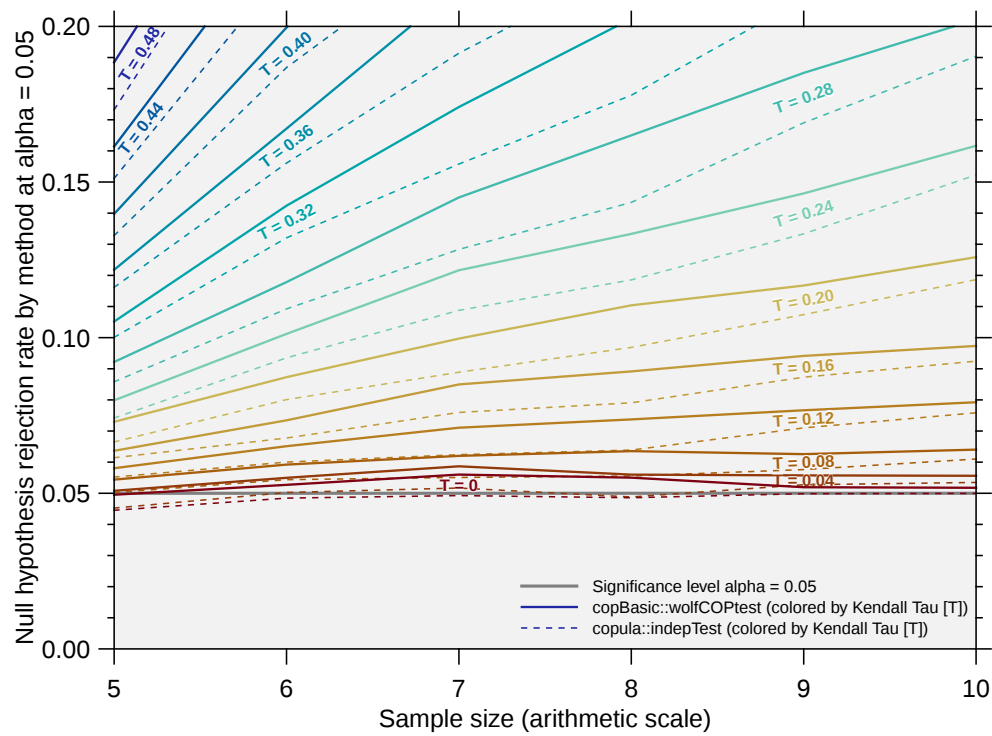


Figure 2. Relation between simulated rejection rates for independence at $\alpha = 0.05$ significance level for selected sample sizes 5–20 for a Clayton copula parameterized by given Kendall Tau as indicated

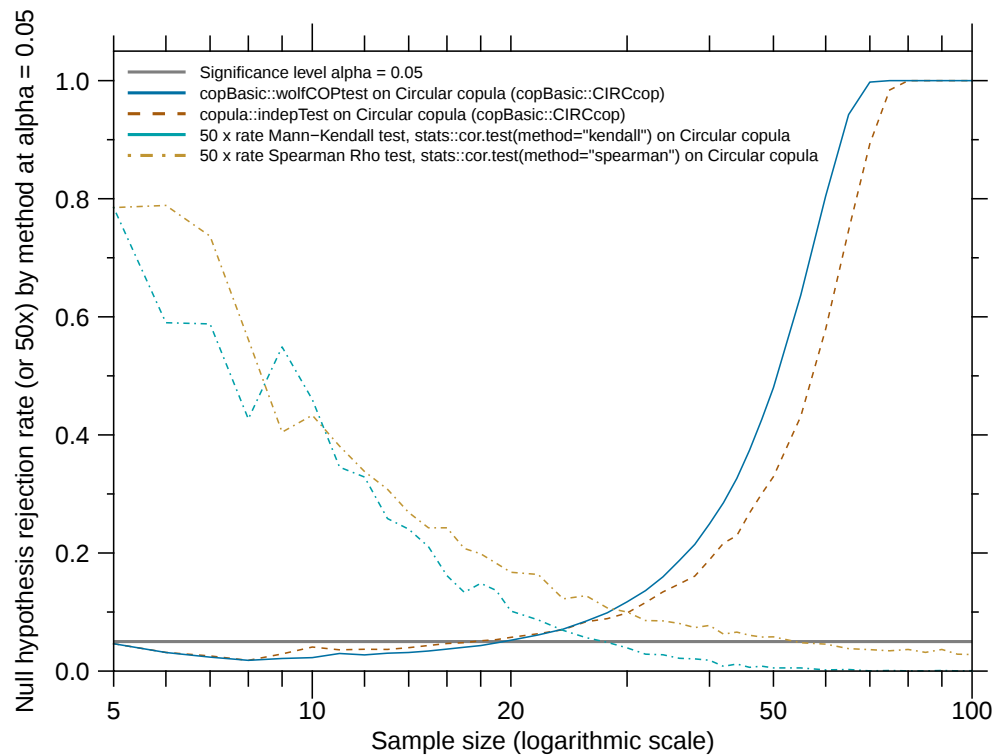


Figure 3. Relation between simulated rejection rates for independence at $\alpha = 0.05$ significance level for selected sample sizes 5–100 for a Circular copula comprised of nothing but singular components

Author(s)

W.H. Asquith

References

- Deheuvels, P., 1981, A non parametric test for independence: Publications de l'Institut de Statistique de l'Université de Paris, v. 26, no. 2, pp. 29–50. [Also available at <https://hal.science/hal-04083801v1>]
- Genest, C. and Rémillard, B., 2004, Tests of independence and randomness based on the empirical copula process: Test, v. 13, pp. 335–369, doi:10.1007/BF02595777.
- Genest, C., Quessy, J.-F., and Rémillard, B., 2007, Asymptotic local efficiency of Cramér–von Mises tests for multivariate independence: The Annals of Statistics, v. 35, pp. 166–191, doi:10.1214/009053606000000984.
- Hofert, M., Kojadinovic, I., Mächler, M., and Yan, J., 2017, Elements of copula modeling with R: Cham, Switzerland, Springer, 267 p., doi:10.1007/9783319896359.
- Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also[wolfCOPtest](#), [CLcop](#)

wolfCOPtest_data_smlsam

*Small Sample Components for Schweizer–Wolff Sigma Test***Description**

Data for the *test* components for small samples. Empirical sampling distributions of *Schweizer–Wolff Sigma* ($\hat{\sigma}_C$, [wolfCOP](#)) for $n \in 3, \dots, 40$ of the NULL : $C = \Pi$ (*Independence copula*, [P](#)), where σ_C is defined under [wolfCOP](#), using simulation sizes of 900,000 for each n . So, total simulation size was 34,200,000. Continuing, this data set is a concatenated table of empirical distributions formed from the unique number of $\hat{\sigma}_C$ (discrete variables in small samples), which large-simulation attainable up to ± 4.26 standard deviations, were computed and tabulated in the `wolfCOPtest_data_smlsam` table. Algorithms for the small-sample components are located at `inst/make_wolfCOPtest/smlsam/make_wolfCOPtestP_smlsam.R`, and this script is the origin of the $n = 40$ in [wolfCOPtest](#) source code.

For the smallest of sample sizes, the *empirical distribution* is a step function, and for each plateau, a nudge using `.Machine$double.eps` has been used to make `approx()` for sensible in [wolfCOPtest](#). In the **Examples** herein, the jaggedness of the empirical distribution can readily be seen and visually smoothing by about $n > 10$. Finally, the [wolfCOPtest](#) function internally loads the `wolfCOPtest_data_smlsam` tables, and most users will not need to access this dataset themselves unless for curiosity.

Usage

```
data(wolfCOPtest_data_smlsam)
```

Format

An R data.frame titled `wolfCOPtest_data_smlsam` with

n Sample size;

probs Nonexceedance probability of empirical distribution of $\hat{\sigma}_C$ ([wolfCOP](#)). Note, `quantile(..., type=4)` was used for computations because of structural similarity to sample $\hat{\sigma}_C$ as implemented in **copBasic**;

wolfemp Value for $\hat{\sigma}_C$ ([wolfCOP](#));

See Also[wolfCOPtest](#)

Examples

```

data(wolfCOPtest_data_smlsam)

## Not run:
# Demonstrate the empirical distributions of Schweizer-Wolff against a parametric
# distribution model. We see quite quickly that the distribution model mimics
# the empirical distribution at quite small sample sizes.
data("wolfCOPtest_data_smlsam")
ff <- pnorm(seq(-3, 3, by=0.1))
plot(c(-3, 3), c(0,1), type="n", xlab="Standard normal variate",
      ylab="Schweizer-Wolff Sigma given C(u,v)=PI")
ns <- c(3, 5, 10, 20, 40)
for(n in ns) {
  sata <- wolfCOPtest_data_smlsam[wolfCOPtest_data_smlsam$n == n,]
  lines(qnorm(sata$probs), sata$wolfemp, col="red", lwd=2)
}
ns <- c(3, 4, 5, 7, 10, 15, 20, 30, 40, 50, 70, 100, 130, 200, 500, 1000, 5000)
for(n in ns) {
  wt <- wolfCOPtest(0, n) # next isolate the distribution type
  # If copy and paste from R help, then one needs to add backslashes to avoid
  # unrecognized escape errors from R in the next line of code.
  type <- gsub("\\\\$", "", gsub("p.value\\(dist_", "", names(wt$p.value)[1]))
  para <- lmomco::vec2par(wt$distpara_by_lmoms, type=type) # type == pe3
  qq <- 1/(1 + exp(-lmomco::qlmomco(ff, para)))
  lines(qnorm(ff), qq, col="blue")
  text(0, 1/(1 + exp(-lmomco::qlmomco(0.5, para))), n, cex=0.8, font=2)
}
par(lheight=0.8)
txt <- c(paste0(lmomco::prettydist(type), " model by estimated L-moments\n",
               "of Sigma distribution for select sample sizes"),
        paste0("Empirical distribution for select sample sizes\n",
               "in wolfCOPtest_data_smlsam data object"))
legend("topleft", txt, box.lty=0, inset=0.01, cex=0.8, adj=c(0, 0.8),
       y.intersp=1.6, lwd=c(1,2), col=c("blue", "red"), bg=NA)
par(lheight=1) #
## End(Not run)

## Not run:
W <- seq(0, 1, by=0.01) # Span all Schweizer-Wolff Sigma, when the first and second
# arguments to wolfCOPtest are scalars, these are treated as sigma and sample size.
for(i in 3:40) { # samples sizes 3 to 40 in wolfCOPtest_data_smlsam.
  Pd <- sapply(W, function(w) wolfCOPtest(w, i, usepade=FALSE)$p.value[1])
  Pe <- sapply(W, function(w) wolfCOPtest(w, i, usepade=FALSE)$p.value[2])
  Pp <- sapply(W, function(w) wolfCOPtest(w, i, usepade=TRUE)$p.value[1])
  # p-value by the generalized normal (parametric) distribution
  plot(W, Pd, type="l", col="grey70", lty=2, lwd=7, main=paste0("n=", i),
        xlab="Schweizer-Wolff Sigma", ylab="Exceedance probability (p-value)")
  points(W, Pp, col="skyblue2", lwd=1) # large sample p-values
  # p-value derived from wolfCOP_data_smlsam tables:
  lines(W, Pe, col="green4", lwd=4) # small sample p-values
} #
## End(Not run)

```

Description

Compute shuffles of *Fréchet–Hoeffding lower-bound copula* (Nelsen, 2006, p. 173), which is defined by partitioning $\mathbf{W}$ within $\mathcal{I}^2$ into n subintervals:

$$\mathbf{W}_n(u, v) = \max\left(\frac{k-1}{n}, u + v - \frac{k}{n}\right)$$

for points within the partitions

$$(u, v) \in \left[\frac{k-1}{n}, \frac{k}{n}\right] \times \left[\frac{k-1}{n}, \frac{k}{n}\right], k = 1, 2, \dots, n$$

and for points otherwise out side the partitions

$$\mathbf{W}_n(u, v) = \min(u, v).$$

The support of $\mathbf{W}_n$ consists of n line segments connecting coordinate pairs $\{(k-1)/n, k/n\}$ and $\{k/n, (k-1)/n\}$ as stated by Nelsen (2006). The *Spearman Rho* ([rhoCOP](#)) is defined by $\rho_C = 1 - (2/n^2)$, and the *Kendall Tau* ([tauCOP](#)) by $\tau_C = 1 - (2/n)$.

Usage

W_N5p12a(u, v, para=1, ...)

Arguments

u	Nonexceedance probability u in the X direction;
v	Nonexceedance probability v in the Y direction;
para	A positive integer $n \in 1, 2, \dots$; and
...	Additional arguments to pass.

Value

Value(s) for the copula are returned.

Author(s)

W.H. Asquith

References

Nelsen, R.B., 2006, An introduction to copulas: New York, Springer, 269 p.

See Also

[W](#), [ORDSUMcop](#), [ORDSUWcop](#), [M_N5p12b](#)

Examples

```
W_N5p12a(0.4, 0.6, para=5)

## Not run:
# Nelsen (2006, exer. 5.12a, p. 172, fig. 5.3a)
UV <- simCOP(1000, cop=W_N5p12a, para=4) # which is the same as
para <- list(cop=c(W, W, W, W), para=NULL, part=c(0,0.25,0.50,0.75,1))
UV <- simCOP(1000, cop=ORDSUMcop, para=para)
## End(Not run)
```

Index

- * **AIC**
 - mleCOP, [265](#)
 - vuongCOP, [366](#)
- * **API to the copula package**
 - GHcop, [147](#)
 - med.regressCOP, [261](#)
 - mleCOP, [265](#)
 - spectralmeas, [333](#)
- * **Akaike Information Criterion**
 - aicCOP, [15](#)
- * **Akaike**
 - mleCOP, [265](#)
 - vuongCOP, [366](#)
- * **Ali–Mikhail–Haq copula**
 - AMHcop, [17](#)
- * **Ali–Mikhail–Haq copula**
 - AMHcop, [17](#)
- * **Asymmetric Gumbel–Hougaard copula**
 - GHcop, [147](#)
- * **BIC**
 - mleCOP, [265](#)
 - vuongCOP, [366](#)
- * **Bekrizade et al. (2012) Examples and Exercises**
 - FGMcop, [134](#)
- * **Bernstein copula**
 - EMPIRcop, [97](#)
 - EMPIRgrid_fast, [113](#)
- * **Bernstein empirical copula**
 - EMPIRcop, [97](#)
 - EMPIRgrid_fast, [113](#)
- * **Bernstein polynomial**
 - EMPIRcop, [97](#)
 - EMPIRgrid_fast, [113](#)
- * **Blomqvist (Schmid-Schmidt) Beta**
 - blomCOPss, [43](#)
 - taildepCOP, [348](#)
- * **Blomqvist Beta**
 - blomatrixCOP, [36](#)
 - blomCOP, [40](#)
 - blomCOPss, [43](#)
- * **C-vine copula**
 - FRcop, [140](#)
- * **C-vines**
 - FRcop, [140](#)
- * **C-vine**
 - FRcop, [140](#)
- * **Clayton copula**
 - CLcop, [52](#)
- * **Conditional L-moments of U given V**
 - LMRuvCOP, [250](#)
- * **Conditional L-moments of V given U**
 - LMRvuCOP, [251](#)
- * **Cuadras–Auge copula**
 - M0cop, [270](#)
- * **Cuadras–Auge copula**
 - M0cop, [270](#)
- * **Debye function**
 - FRcop, [140](#)
- * **Durante and Sempi (2015) Examples and Exercises**
 - kfuncCOP, [203](#)
- * **Erdely (2017) Examples and Exercises**
 - glueCOP, [160](#)
- * **Farlie–Gumbel–Morgenstern copula**
 - FGMcop, [134](#)
- * **Farlie–Gumbel–Morgenstern copula**
 - FGMcop, [134](#)
- * **Footrule**
 - footCOP, [137](#)
- * **Frank copula**
 - FRcop, [140](#)
- * **Frechet Family copula**
 - FRECHETcop, [143](#)
- * **Frechet lower bound**
 - W, [372](#)
- * **Frechet lower-bound copula**
 - W, [372](#)

- * **Frechet upper bounds**
M, 260
- * **Frechet upper-bound copula**
M, 260
- * **Frechet–Hoeffding lower-bound copula**
W, 372
- * **Frechet–Hoeffding lower-bound**
W, 372
- * **Frechet–Hoeffding upper bounds**
M, 260
- * **Frechet–Hoeffding upper-bound copula**
M, 260
- * **Frechet–Hoeffding lower-bound copula**
W, 372
- * **Frechet–Hoeffding lower-bound**
W, 372
- * **Frechet–Hoeffding upper bounds**
M, 260
- * **Frechet–Hoeffding upper-bound copula**
M, 260
- * **GNO (see generalized normal distribution)**
wolfCOPtest, 384
- * **Galambos copula (lower extreme value form)**
GLcop, 157
- * **Galambos copula**
GLcop, 157
- * **Galambos extreme value copula**
GLcop, 157
- * **Gaussian copula**
blomCOP, 40
med.regressCOP, 261
NORMcop, 276
- * **Gumbel–Hougaard copula**
GHcop, 147
- * **Gumbel–Hougaard extreme value copula**
GHcop, 147
- * **Gumbel–Hougaard extreme value copula**
GHcop, 147
- * **Hairpin copula**
HAIRPINcop, 164
- * **Halton sequence (Monte Carlo integration)**
bilmoms, 30
- * **Hazen copula**
EMPIRCop, 97
- * **Hazen empirical copula**
EMPIRCop, 97
- * **Hoeffding Phi-Square**
hoefCOP, 165
- * **Hoeffding Phi**
hoefCOP, 165
- * **Hofert et al. (2018) Examples and Exercises**
composite1COP, 55
- * **Husler–Reiss copula**
HRcop, 169
- * **Husler–Reiss extreme value copula**
HRcop, 169
- * **Husler–Reiss copula**
HRcop, 169
- * **Husler–Reiss extreme value copula**
HRcop, 169
- * **Hypothesis test for Schweizer and Wolff Sigma**
wolfCOPtest, 384
wolfCOPtest_check, 391
- * **I-divergence**
kullCOP, 218
- * **Jeffery Divergence**
kullCOP, 218
- * **Jeffery’s Divergence**
kullCOP, 218
- * **Joe (2014) Examples and Exercises**
blomCOP, 40
CLcop, 52
densityCOP, 76
densityCOPplot, 78
derCOPinv, 85
FRcop, 140
GHcop, 147
GLcop, 157
HRcop, 169
JOcopB5, 182
joeskewCOP, 184
kfuncCOP, 203
kullCOP, 218
PLACKETTcop, 289
rhoCOP, 313
semicorCOP, 320
tailordCOP, 353
tEVcop, 361
- * **Joe–Ma copula**
JOMAcop, 200
- * **Joe/B5 copula**
JOcopB5, 182

- * **Joe/BB4 copula**
GLcop, 157
- * **Kendall Function (L-moments)**
kfuncCOPlmoms, 213
- * **Kendall Function L-moment diagram**
kfuncCOPlmoms, 213
- * **Kendall Function L-moment ratio diagram**
kfuncCOPlmoms, 213
- * **Kendall Function L-moments**
kfuncCOPlmoms, 213
- * **Kendall Function Lmoment diagram**
kfuncCOPlmoms, 213
- * **Kendall Function Lmoment ratio diagram**
kfuncCOPlmoms, 213
- * **Kendall Function Lmoments**
kfuncCOPlmoms, 213
- * **Kendall Function**
kfuncCOP, 203
kfuncCOP_Pd, 216
W, 372
- * **Kendall Measure**
kfuncCOP, 203
- * **Kendall Tau**
tauCOP, 355
- * **Kendall distribution function**
kfuncCOP, 203
W, 372
- * **Kendall function**
kfuncCOP, 203
W, 372
- * **Kendall inverse distribution function**
kfuncCOPinv, 211
- * **Kendall measure**
kfuncCOP, 203
W, 372
- * **Kendall quantile function**
kfuncCOPinv, 211
- * **Khoudraji device (with independence)**
composite1COP, 55
- * **Khoudraji device**
composite1COP, 55
composite2COP, 57
- * **Kullback–Leibler Divergence**
kullCOP, 218
- * **Kullback–Leibler sample size**
kullCOP, 218
- * **Kullback–Leibler**
mleCOP, 265
vuongCOP, 366
- * **Kullback-Leibler Divergence**
kullCOP, 218
- * **Kullback-Leibler sample size**
kullCOP, 218
- * **Kullback-Leibler**
mleCOP, 265
vuongCOP, 366
- * **L-comoment copula inference**
vuongCOP, 366
- * **L-comoment ratio diagram**
FGMcop, 134
lcomCOP, 222
LCOMDIA_GH2cop, 229
LCOMDIA_GH3cop, 232
LCOMDIA_ManyCops, 233
ORDSUMcop, 278
- * **L-comoments (ratio diagram)**
FGMcop, 134
lcomCOP, 222
LCOMDIA_GH2cop, 229
LCOMDIA_GH3cop, 232
LCOMDIA_ManyCops, 233
ORDSUMcop, 278
- * **L-comoments**
bilmoms, 30
copBasic-package, 5
isCOP.permsym, 172
isCOP.radsym, 176
lcomCOP, 222
lcomCOPpv, 226
lcomoms2.ABcop2parameter, 238
lcomoms2.ABKGcop2parameter, 242
M0cop, 270
P, 286
simcomposite3COP, 323
simcompositeCOP, 325
- * **L-moments (conditional) of U given V**
LMRuvCOP, 250
- * **L-moments (conditional) of V given U**
LMRvuCOP, 251
- * **L-moments by CDF**
kfuncCOPlmoms, 213
- * **L-moments of combined U and V**
uvlmoms, 364
- * **L-moments**
kfuncCOPlmoms, 213

- uvlmoms, 364
- * **Lcomoment copula inference**
 - vuongCOP, 366
- * **Lcomoments**
 - bilmoms, 30
 - copBasic-package, 5
 - isCOP.permsym, 172
 - isCOP.radsym, 176
 - lcomCOP, 222
 - MOcop, 270
 - P, 286
- * **Linear Spearman copula**
 - FRECHETcop, 143
- * **Linear Spearman**
 - FRECHETcop, 143
- * **Lmoments**
 - kfuncCOPlmoms, 213
 - uvlmoms, 364
- * **Lp distance**
 - hoefCOP, 165
- * **M-ordinal sum of the summands**
 - ORDSUMcop, 278
- * **Markov Process**
 - prod2COP, 294
- * **Markov Product**
 - prod2COP, 294
- * **Markov process**
 - prod2COP, 294
- * **Markov product**
 - prod2COP, 294
- * **Marshall–Olkin copula**
 - MOcop, 270
- * **Marshall–Olkin copula**
 - MOcop, 270
- * **Moments of combined U and V**
 - joeskewCOP, 184
- * **Monte Carlo integration**
 - wolfCOPsamc, 379
- * **Nelsen (2006) Examples and Exercises**
 - asCOP, 20
 - bilmoms, 30
 - blomCOP, 40
 - CIRCcop, 49
 - copBasic-package, 5
 - derCOPinv, 85
 - hoefCOP, 165
 - isfuncCOP, 181
 - M_N5p12b, 272
 - prod2COP, 294
 - RFcop, 309
 - taildepCOP, 348
 - tauCOP, 355
 - TRICop, 363
 - W_N5p12a, 399
- * **Normal copula**
 - blomCOP, 40
 - med.regressCOP, 261
 - NORMcop, 276
- * **Nu-Skew and Nu-Star (skewnesses)**
 - joeskewCOP, 184
- * **Ordinal Sums**
 - ORDSUMcop, 278
 - ORDSUWcop, 284
- * **PE3 (see Pearson type III distribution)**
 - wolfCOPtest, 384
 - wolfCOPtest_data_smlsam, 397
- * **Package Lmoments**
 - wolfCOPsamc, 379
- * **Package copula**
 - wolfCOPtest, 384
 - wolfCOPtest_check, 391
- * **Package ellipse**
 - wolfCOPsamc, 379
- * **Package parallel**
 - wolfCOPsamc, 379
- * **Package randtoolbox**
 - bilmoms, 30
 - kullCOP, 218
 - wolfCOPsamc, 379
- * **Pareto copula**
 - PARETOcop, 287
- * **Pearson correlation coefficient**
 - gEVcop, 145
 - P, 286
 - rhoCOP, 313
- * **Pearson type III distribution**
 - wolfCOPtest, 384
 - wolfCOPtest_data_smlsam, 397
- * **Plackett copula**
 - PLACKETTcop, 289
 - PLACKETTpar, 291
 - PLACKETTsim, 293
- * **Raftery copula**
 - RFcop, 309
- * **Rayleigh copula**
 - RAYcop, 305

- * **Rosenblatt transform**
simCOP, [327](#)
- * **Salvadori et al. (2007) Examples and Exercises**
copBasic-package, [5](#)
derCOPinv, [85](#)
kfuncCOP, [203](#)
surfuncCOP, [345](#)
- * **Schweizer and Wolff Sigma (hypothesis test)**
wolfCOPtest, [384](#)
wolfCOPtest_check, [391](#)
- * **Schweizer and Wolff Sigma**
wolfCOP, [374](#)
wolfCOPsamc, [379](#)
wolfCOPtest, [384](#)
wolfCOPtest_check, [391](#)
wolfCOPtest_data_smlsam, [397](#)
- * **Shuffle of Frechet lower bound**
W_N5p12a, [399](#)
- * **Shuffle of Frechet lower-bound copula**
W_N5p12a, [399](#)
- * **Shuffle of Frechet upper bounds**
M_N5p12b, [272](#)
- * **Shuffle of Frechet upper-bound copula**
M_N5p12b, [272](#)
- * **Shuffle of Frechet–Hoeffding lower-bound copula**
W_N5p12a, [399](#)
- * **Shuffle of Frechet–Hoeffding lower-bound**
W_N5p12a, [399](#)
- * **Shuffle of Frechet–Hoeffding upper bounds**
M_N5p12b, [272](#)
- * **Shuffle of Frechet–Hoeffding upper-bound copula**
M_N5p12b, [272](#)
- * **Shuffle of Frechet-Hoeffding lower-bound copula**
W_N5p12a, [399](#)
- * **Shuffle of Frechet-Hoeffding lower-bound**
W_N5p12a, [399](#)
- * **Shuffle of Frechet-Hoeffding upper bounds**
M_N5p12b, [272](#)
- * **Shuffle of Frechet-Hoeffding upper-bound copula**
M_N5p12b, [272](#)
- * **Sobol sequence (Monte Carlo integration)**
kullCOP, [218](#)
- * **Spearman Footrule**
footCOP, [137](#)
- * **Spearman Rho**
bilmoms, [30](#)
gEvcop, [145](#)
lcomCOP, [222](#)
rhoCOP, [313](#)
- * **Student t**
Tcop, [358](#)
tEvcop, [361](#)
- * **Table of Copulas**
copBasic-package, [5](#)
- * **Table of Expectations**
copBasic-package, [5](#)
- * **Table of Probabilities and Expectations**
copBasic-package, [5](#)
- * **Table of Probabilities**
copBasic-package, [5](#)
- * **Theory Examples (reflection/rotation)**
COP, [65](#)
EuvCOP, [129](#)
EvuCOP, [131](#)
JOcopB5, [182](#)
LzCOPpermsym, [254](#)
RFcop, [309](#)
semicorCOP, [320](#)
simCOPmicro, [329](#)
- * **Triangular copula**
TRICop, [363](#)
- * **Vine Copula**
FRcop, [140](#)
- * **Vuong Procedure**
mleCOP, [265](#)
vuongCOP, [366](#)
- * **Vuong**
mleCOP, [265](#)
vuongCOP, [366](#)
- * **W-ordinal sum of the summands**
ORDSUWcop, [284](#)
- * **Weibull copula**
EMPIRcop, [97](#)
- * **Weibull empirical copula**
EMPIRcop, [97](#)
- * **Zhang and Singh (2019) Examples and Exercises**
derCOPinv, [85](#)

- * **artificial bee optimization**
 - LzCOPpermsym, 254
- * **asymmetric Gumbel copula**
 - GHcop, 147
- * **asymmetric Gumbel–Hougaard copula**
 - GHcop, 147
- * **asymmetric Gumbel–Hougaard copula**
 - GHcop, 147
- * **asymmetric logistic copula**
 - GHcop, 147
- * **bivariate L-correlation**
 - bilmoms, 30
 - lcomCOP, 222
- * **bivariate L-kurtosis**
 - bilmoms, 30
 - lcomCOP, 222
- * **bivariate L-moments**
 - bilmoms, 30
 - lcomCOP, 222
- * **bivariate L-skew**
 - bilmoms, 30
 - lcomCOP, 222
- * **bivariate asymmetry (measures)**
 - bilmoms, 30
 - lcomCOP, 222
 - uvlmoms, 364
- * **bivariate prediction**
 - bicoploc, 24
- * **bivariate skewness**
 - bilmoms, 30
 - joeskewCOP, 184
 - lcomCOP, 222
 - uvlmoms, 364
- * **censored data**
 - wolfCOPtest, 384
- * **circular copula**
 - CIRCcop, 49
- * **comprehensive copula**
 - FRcop, 140
 - FRECHETcop, 143
 - JOMAcop, 200
 - PLACKETTcop, 289
- * **concordance function**
 - tauCOP, 355
- * **conditional approach**
 - simCOP, 327
- * **conditional cumulative distribution function**
 - derCOP, 81
 - derCOP2, 83
- * **conditional expectation**
 - EuvCOP, 129
 - EvuCOP, 131
 - LMRuvCOP, 250
 - LMRvuCOP, 251
- * **conditional quantile function**
 - derCOPinv, 85
 - derCOPinv2, 89
- * **conditional return period**
 - surfuncCOP, 345
- * **copula (characteristics)**
 - blomatrixCOP, 36
 - blomCOP, 40
 - blomCOPss, 43
 - diagCOP, 90
 - diagCOPatf, 92
 - footCOP, 137
 - giniCOP, 154
 - hoefCOP, 165
 - isCOP.LTD, 171
 - isCOP.permsym, 172
 - isCOP.PQD, 173
 - isCOP.radsym, 176
 - isCOP.RTI, 180
 - joeskewCOP, 184
 - kfuncCOP, 203
 - kfuncCOP_Pd, 216
 - kfuncCOPinv, 211
 - kfuncCOPlmoms, 213
 - lcomCOPpv, 226
 - LzCOPpermsym, 254
 - rhobevCOP, 311
 - rhoCOP, 313
 - semicorCOP, 320
 - tailconCOP, 347
 - taildepCOP, 348
 - tailordCOP, 353
 - tauCOP, 355
 - wolfCOP, 374
 - wolfCOPsamc, 379
- * **copula (comprehensive)**
 - FRcop, 140
 - FRECHETcop, 143
 - JOMAcop, 200
 - PLACKETTcop, 289
- * **copula (conditional distribution)**

- COP, 65
- derCOP, 81
- derCOP2, 83
- EuvCOP, 129
- EvuCOP, 131
- LMRuvCOP, 250
- LMRvuCOP, 251
- simCOPmicro, 329
- * **copula (conditional quantile function)**
 - COP, 65
 - derCOPinv, 85
 - derCOPinv2, 89
 - simCOPmicro, 329
- * **copula (density)**
 - densityCOP, 76
 - densityCOPplot, 78
 - kullCOP, 218
 - PLACKETTcop, 289
 - vuongCOP, 366
- * **copula (derivative inverse)**
 - derCOPinv, 85
 - derCOPinv2, 89
- * **copula (derivative)**
 - derCOP, 81
 - derCOP2, 83
- * **copula (diagonal inversion)**
 - bicoploc, 24
 - diagCOPatf, 92
- * **copula (diagonal)**
 - bicoploc, 24
 - diagCOP, 90
 - diagCOPatf, 92
- * **copula (empirical)**
 - EMPIRcop, 97
- * **copula (estimation)**
 - PLACKETTpar, 291
- * **copula (extreme value)**
 - gEVcop, 145
 - GHcop, 147
 - GLcop, 157
 - HRcop, 169
 - tEVcop, 361
- * **copula (fitting) Metaheuristic Optimization**
 - LzCOPpermsym, 254
- * **copula (fitting)**
 - copBasic.fitpara, 70
 - LzCOPpermsym, 254
 - mleCOP, 265
- * **copula (formulas)**
 - AMHcop, 17
 - asCOP, 20
 - CIRCcop, 49
 - CLcop, 52
 - COP, 65
 - FGMcop, 134
 - FRcop, 140
 - FRECHETcop, 143
 - gEVcop, 145
 - GHcop, 147
 - GLcop, 157
 - glueCOP, 160
 - HAIRPINcop, 164
 - HRcop, 169
 - JOcopB5, 182
 - JOMAcop, 200
 - M, 260
 - M_N5p12b, 272
 - M0cop, 270
 - N4212cop, 274
 - N4220cop, 275
 - NORMcop, 276
 - ORDSUMcop, 278
 - ORDSUWcop, 284
 - P, 286
 - PARET0cop, 287
 - PLACKETTcop, 289
 - PSP, 299
 - RAYcop, 305
 - RFcop, 309
 - rN4220cop, 317
 - Tcop, 358
 - tEVcop, 361
 - TRIcop, 363
 - W, 372
 - W_N5p12a, 399
- * **copula (goodness-of-fit)**
 - aicCOP, 15
 - bicCOP, 22
 - copBasic.fitpara, 70
 - psepolar, 297
 - rmseCOP, 316
 - spectralmeas, 333
 - statTn, 340
 - vuongCOP, 366
- * **copula (hypothesis test)**

- wolfCOPtest, 384
- wolfCOPtest_check, 391
- * **copula (inference)**
 - aicCOP, 15
 - bicCOP, 22
 - copBasic.fitpara, 70
 - kullCOP, 218
 - lcomCOPpv, 226
 - psepolar, 297
 - rmseCOP, 316
 - spectralmeas, 333
 - statTn, 340
 - vuongCOP, 366
- * **copula (inverse)**
 - COPinv, 73
 - COPinv2, 74
- * **copula (limits/bounds)**
 - M, 260
 - M_N5p12b, 272
 - W, 372
 - W_N5p12a, 399
- * **copula (properties for extreme value)**
 - rhobevCOP, 311
- * **copula (properties)**
 - blomatrixCOP, 36
 - blomCOP, 40
 - blomCOPss, 43
 - footCOP, 137
 - giniCOP, 154
 - hoefCOP, 165
 - isCOP.LTD, 171
 - isCOP.permsym, 172
 - isCOP.PQD, 173
 - isCOP.radsym, 176
 - isCOP.RTI, 180
 - isfuncCOP, 181
 - joeskewCOP, 184
 - kfuncCOP, 203
 - kfuncCOP_Pd, 216
 - kfuncCOPinv, 211
 - LzCOPpermsym, 254
 - rhobevCOP, 311
 - rhoCOP, 313
 - semicorCOP, 320
 - tailconCOP, 347
 - taildepCOP, 348
 - tailordCOP, 353
 - tauCOP, 355
 - wolfCOP, 374
 - wolfCOPsamc, 379
- * **copula (reflection)**
 - COP, 65
 - surCOP, 343
- * **copula (rotation)**
 - COP, 65
- * **copula (simulation)**
 - PLACKETTsim, 293
 - simcomposite3COP, 323
 - simcompositeCOP, 325
 - simCOP, 327
 - simCOPmicro, 329
- * **copula (symmetry)**
 - isCOP.permsym, 172
 - isCOP.radsym, 176
 - LzCOPpermsym, 254
- * **copula (tail characteristics)**
 - blomCOPss, 43
 - semicorCOP, 320
 - tailconCOP, 347
 - taildepCOP, 348
 - tailordCOP, 353
- * **copula (tail properties)**
 - blomCOPss, 43
 - semicorCOP, 320
 - tailconCOP, 347
 - taildepCOP, 348
 - tailordCOP, 353
- * **copula (utilities)**
 - asCOP, 20
- * **copula (utility)**
 - COPinv, 73
 - COPinv2, 74
 - derCOP, 81
 - derCOP2, 83
 - derCOPinv, 85
 - derCOPinv2, 89
 - EuvCOP, 129
 - EvuCOP, 131
 - LMRuvCOP, 250
 - LMRvuCOP, 251
- * **copula L-cokurtosis**
 - bilmoms, 30
 - lcomCOP, 222
- * **copula L-comoments**
 - bilmoms, 30
 - lcomCOP, 222

- * **copula L-correlation**
 - bilmoms, [30](#)
 - lcomCOP, [222](#)
- * **copula L-coskew**
 - bilmoms, [30](#)
 - lcomCOP, [222](#)
- * **copula composition (convex combination)**
 - convex2COP, [62](#)
 - convexCOP, [64](#)
- * **copula composition (four compositing parameters)**
 - composite3COP, [60](#)
- * **copula composition (permutation asymmetric)**
 - breveCOP, [46](#)
- * **copula composition (two compositing parameters with independence)**
 - composite1COP, [55](#)
- * **copula composition (two compositing parameters)**
 - composite1COP, [55](#)
 - composite2COP, [57](#)
- * **copula composition**
 - breveCOP, [46](#)
 - composite1COP, [55](#)
 - composite2COP, [57](#)
 - composite3COP, [60](#)
 - convex2COP, [62](#)
 - convexCOP, [64](#)
 - glueCOP, [160](#)
 - lcomoms2.ABcop2parameter, [238](#)
 - lcomoms2.ABKGcop2parameter, [242](#)
 - simcomposite3COP, [323](#)
 - simcompositeCOP, [325](#)
- * **copula covariance**
 - rhoCOP, [313](#)
- * **copula divergence**
 - kullCOP, [218](#)
- * **copula gluing**
 - glueCOP, [160](#)
- * **copula multiplication**
 - prod2COP, [294](#)
- * **copula operator**
 - coCOP, [53](#)
 - COP, [65](#)
 - COPinv, [73](#)
 - COPinv2, [74](#)
 - derCOP, [81](#)
 - derCOP2, [83](#)
 - derCOPinv, [85](#)
 - derCOPinv2, [89](#)
 - duCOP, [94](#)
 - EuvCOP, [129](#)
 - EvuCOP, [131](#)
 - LMRuvCOP, [250](#)
 - LMRvuCOP, [251](#)
 - prod2COP, [294](#)
 - surCOP, [343](#)
 - surfuncCOP, [345](#)
- * **copula product**
 - prod2COP, [294](#)
- * **copula section**
 - sectionCOP, [318](#)
- * **copula skew**
 - joeskewCOP, [184](#)
- * **copula theory**
 - copBasic-package, [5](#)
- * **copula utility**
 - gridCOP, [162](#)
- * **copulatic surface**
 - EMPIRgrid, [103](#)
 - EMPIRgrid_fast, [113](#)
 - EMPIRgrid_lkup, [115](#)
 - EMPIRgridder, [107](#)
 - EMPIRgridder2, [109](#)
 - EMPIRgridderinv, [110](#)
 - EMPIRgridderinv2, [112](#)
 - gridCOP, [162](#)
- * **copula**
 - AMHcop, [17](#)
 - CIRCcop, [49](#)
 - CLcop, [52](#)
 - EMPIRcop, [97](#)
 - FGMcop, [134](#)
 - FRcop, [140](#)
 - FRECHETcop, [143](#)
 - gEVcop, [145](#)
 - GHcop, [147](#)
 - GLcop, [157](#)
 - HAIRPINcop, [164](#)
 - HRcop, [169](#)
 - JOcopB5, [182](#)
 - JOMAcop, [200](#)
 - M, [260](#)
 - M_N5p12b, [272](#)
 - M0cop, [270](#)

- N4212cop, 274
- N4220cop, 275
- NORMcop, 276
- ORDSUMcop, 278
- ORDSUWcop, 284
- P, 286
- PARETOcop, 287
- PLACKETTcop, 289
- PSP, 299
- RAYcop, 305
- RFcop, 309
- rN4220cop, 317
- Tcop, 358
- tEVcop, 361
- TRICop, 363
- W, 372
- W_N5p12a, 399
- * **cumulative distribution**
 - kfuncCOPlmoms, 213
- * **datasets**
 - ReineckeWell1266, 307
 - ReineckeWells, 308
 - wolfCOPtest_data_smlsam, 397
- * **decomposition of Kendall Tau**
 - kfuncCOP, 203
- * **dependence index**
 - hoefCOP, 165
- * **derivative**
 - derCOP, 81
 - derCOP2, 83
 - derCOPinv, 85
 - derCOPinv2, 89
 - EMPIRgridder, 107
 - EMPIRgridder2, 109
 - EMPIRgridderinv, 110
 - EMPIRgridderinv2, 112
- * **diagnostics**
 - diagCOP, 90
 - diagCOPatf, 92
 - level.curvesCOP, 245
 - level.curvesCOP2, 247
 - level.setCOP, 248
 - level.setCOP2, 249
 - sectionCOP, 318
- * **diagonal inversion**
 - diagCOPatf, 92
- * **empirical copula (derivative inverses)**
 - EMPIRgridderinv, 110
 - EMPIRgridderinv2, 112
- * **empirical copula (derivatives)**
 - EMPIRgridder, 107
 - EMPIRgridder2, 109
- * **empirical copula (median regression)**
 - EMPIRmed.regress, 116
 - EMPIRmed.regress2, 117
- * **empirical copula (quantile regression)**
 - EMPIRmed.regress, 116
 - EMPIRmed.regress2, 117
 - EMPIRqua.regress, 118
 - EMPIRqua.regress2, 124
- * **empirical copula (simulation)**
 - EMPIRsim, 125
 - EMPIRsimv, 127
- * **empirical copula (utility)**
 - EMPIRcopdf, 102
 - EMPIRgrid, 103
 - EMPIRgrid_fast, 113
 - EMPIRgrid_lkup, 115
- * **empirical copula**
 - EMPIRcop, 97
 - EMPIRcopdf, 102
 - EMPIRgrid, 103
 - EMPIRgrid_fast, 113
 - EMPIRgrid_lkup, 115
 - EMPIRgridder, 107
 - EMPIRgridder2, 109
 - EMPIRgridderinv, 110
 - EMPIRgridderinv2, 112
 - EMPIRmed.regress, 116
 - EMPIRmed.regress2, 117
 - EMPIRqua.regress, 118
 - EMPIRqua.regress2, 124
 - EMPIRsim, 125
 - EMPIRsimv, 127
- * **exclusive or**
 - duCOP, 94
- * **expectation**
 - EuvCOP, 129
 - EvuCOP, 131
- * **extended delta index**
 - wolfCOP, 374
- * **extreme value copula**
 - gEVcop, 145
 - GHcop, 147
 - GLcop, 157
 - HRcop, 169

- tEvcop, 361
- * **g-EV copula**
 - gEvcop, 145
- * **generalized Farlie-Gumbel-Morgenstern copula**
 - FGMcop, 134
- * **generalized normal distribution**
 - wolfCOPtest, 384
- * **goodness-of-fit**
 - aicCOP, 15
 - bicCOP, 22
 - copBasic.fitpara, 70
 - psepolar, 297
 - rmseCOP, 316
 - spectralmeas, 333
 - statTn, 340
 - vuongCOP, 366
- * **gridded empirical copula**
 - EMPIRgrid, 103
 - EMPIRgrid_fast, 113
 - EMPIRgrid_lkup, 115
 - EMPIRgridder, 107
 - EMPIRgridder2, 109
 - EMPIRgridderinv, 110
 - EMPIRgridderinv2, 112
- * **hypothesis testing**
 - lcomCOPpv, 226
- * **independence copula**
 - P, 286
- * **independence test**
 - wolfCOPtest, 384
 - wolfCOPtest_check, 391
- * **inference**
 - aicCOP, 15
 - bicCOP, 22
 - copBasic.fitpara, 70
 - kullCOP, 218
 - lcomCOPpv, 226
 - psepolar, 297
 - rmseCOP, 316
 - spectralmeas, 333
 - statTn, 340
 - vuongCOP, 366
- * **iterated Farlie-Gumbel-Morgenstern copula**
 - FGMcop, 134
- * **joint probability (exclusive or)**
 - duCOP, 94
- * **joint probability**
 - coCOP, 53
 - COP, 65
 - duCOP, 94
 - joint.curvesCOP, 189
 - joint.curvesCOP2, 193
 - jointCOP, 195
 - surCOP, 343
 - surfuncCOP, 345
- * **level contours**
 - level.curvesCOP, 245
 - level.curvesCOP2, 247
 - level.setCOP, 248
 - level.setCOP2, 249
- * **level contour**
 - level.curvesCOP, 245
 - level.curvesCOP2, 247
 - level.setCOP, 248
 - level.setCOP2, 249
- * **level curves**
 - level.curvesCOP, 245
 - level.curvesCOP2, 247
 - level.setCOP, 248
 - level.setCOP2, 249
- * **level curve**
 - level.curvesCOP, 245
 - level.curvesCOP2, 247
 - level.setCOP, 248
 - level.setCOP2, 249
- * **level sets**
 - level.curvesCOP, 245
 - level.curvesCOP2, 247
 - level.setCOP, 248
 - level.setCOP2, 249
- * **level set**
 - level.curvesCOP, 245
 - level.curvesCOP2, 247
 - level.setCOP, 248
 - level.setCOP2, 249
- * **line of organic correlation**
 - bicoploc, 24
- * **literature errors and inconsistencies**
 - GHcop, 147
 - kfuncCOP, 203
 - stabtaildepf, 337
 - tauCOP, 355
- * **logit transform**
 - wolfCOPtest, 384

- wolfCOPtest_data_smlsam, 397
- * **maximum likelihood**
 - mleCOP, 265
- * **mean U given V**
 - EuvCOP, 129
- * **mean V given U**
 - EvuCOP, 131
- * **medial correlation coefficient**
 - blomCOP, 40
- * **medial correlation**
 - blomCOP, 40
- * **median regression**
 - med.regressCOP, 261
 - med.regressCOP2, 264
- * **multivariate**
 - copBasic-package, 5
- * **mutually exclusive or condition**
 - duCOP, 94
- * **mutually exclusive or**
 - duCOP, 94
- * **ordinal sums**
 - ORDSUMcop, 278
 - ORDSUWcop, 284
- * **organic correlation**
 - bicoploc, 24
- * **package ABCoptim (application of)**
 - LzCOPpermsym, 254
- * **package acopula (application of)**
 - GHcop, 147
- * **package copula (comparison to)**
 - AMHcop, 17
 - EMPIRCop, 97
 - gEVcop, 145
 - GHcop, 147
 - med.regressCOP, 261
 - mleCOP, 265
 - spectralmeas, 333
 - taildepCOP, 348
 - tEVcop, 361
- * **package gumbel (application of)**
 - GHcop, 147
- * **package lmomco (application of)**
 - P, 286
- * **package metaheuristicOpt (application of)**
 - LzCOPpermsym, 254
- * **package pso (application of)**
 - LzCOPpermsym, 254
- * **package randtoolbox (application of)**
 - kullCOP, 218
 - LzCOPpermsym, 254
- * **particle swarm optimization**
 - LzCOPpermsym, 254
- * **permutation asymmetry**
 - breveCOP, 46
 - joeskewCOP, 184
- * **polar**
 - psepolar, 297
 - spectralmeas, 333
- * **product copula**
 - P, 286
- * **pseudo-polar representation**
 - psepolar, 297
- * **quantile regression**
 - med.regressCOP, 261
 - med.regressCOP2, 264
 - qua.regressCOP, 300
 - qua.regressCOP.draw, 302
 - qua.regressCOP2, 304
- * **radial asymmetry**
 - joeskewCOP, 184
- * **reduced major axis**
 - bicoploc, 24
- * **relative entropy**
 - kullCOP, 218
- * **return period (conditional)**
 - copBasic-package, 5
- * **return period (secondary)**
 - copBasic-package, 5
 - kfuncCOP, 203
- * **semi-correlation coefficient**
 - semicorCOP, 320
- * **semi-correlation**
 - semicorCOP, 320
- * **shuffle (in Examples)**
 - asCOP, 20
- * **shuffle**
 - asCOP, 20
 - M_N5p12b, 272
 - W_N5p12a, 399
- * **spectral measure**
 - spectralmeas, 333
- * **stable tail dependence function**
 - stabtaildepf, 337
- * **summands**
 - ORDSUMcop, 278
 - ORDSUWcop, 284

- * **symmetric Gumbel copula**
 - GHcop, 147
 - * **symmetric Gumbel–Hougaard copula**
 - GHcop, 147
 - * **symmetric Gumbel–Hougaard copula**
 - GHcop, 147
 - * **t-EV copula**
 - tEVcop, 361
 - * **t-Student copula**
 - Tcop, 358
 - * **tail dependence**
 - blomCOPss, 43
 - stabtaildepf, 337
 - taildepCOP, 348
 - * **triangle (in Examples)**
 - asCOP, 20
 - * **visualization**
 - densityCOP, 76
 - densityCOPplot, 78
 - diagCOP, 90
 - diagCOPatf, 92
 - level.curvesCOP, 245
 - level.curvesCOP2, 247
 - level.setCOP, 248
 - level.setCOP2, 249
 - qua.regressCOP.draw, 302
 - sectionCOP, 318
 - simcomposite3COP, 323
 - simcompositeCOP, 325
 - simCOP, 327
- aicCOP, 8, 15, 22, 23, 258, 267, 316, 317, 340, 343
- AMHcop, 6, 17
- asCOP, 20, 32, 187
- bicCOP, 8, 16, 22, 258, 267, 316, 317, 340, 343
- bicoploc, 24
- bilmoms, 7, 8, 30, 187, 224, 225, 227, 365
- blomatrixCOP, 36, 45
- blomatrixCOPdec, 41
- blomatrixCOPdec (blomatrixCOP), 36
- blomatrixCOPiqr, 41
- blomatrixCOPiqr (blomatrixCOP), 36
- blomCOP, 7, 36, 37, 40, 43–45, 99, 100, 139, 155, 168, 173, 175, 186, 188, 263, 314, 342, 347, 351, 357, 377
- blomCOPss, 36, 37, 40, 41, 43, 347, 350, 351
- breveCOP, 11, 46, 56, 59, 61, 63, 65, 161, 210, 270, 373
- CIRCcop, 6, 49, 393
- CLcop, 5, 6, 52, 256, 391, 392, 397
- coCOP, 7, 10, 12, 53, 68, 94, 96, 192, 344
- composite1COP, 9, 11, 46, 47, 55, 56, 58, 59, 61, 63, 65, 66, 68, 71, 161, 256, 294, 296, 323, 325
- composite2COP, 11, 46, 47, 55, 56, 57, 60, 61, 63, 65, 66, 161, 163, 191, 240, 294, 296, 323, 325
- composite3COP, 11, 46, 47, 56, 58, 59, 60, 63, 65, 66, 161, 244, 294, 296, 323
- concordCOP, 137, 138, 154
- concordCOP (tauCOP), 355
- convex2COP, 11, 47, 62, 65, 66, 144, 154, 294, 296
- convexCOP, 11, 47, 56, 59, 61–63, 64, 66, 161, 294, 296
- COP, 7, 10, 12, 13, 21, 32, 43, 47, 54–56, 59, 61, 63, 65, 65, 74, 75, 91, 96, 161, 164, 176, 197, 228, 275, 296, 309, 319, 322, 329, 331, 343, 344, 347, 351, 354, 366
- copBasic-package, 5
- copBasic.fitpara, 70
- copBasic.fitpara.beta, 341
- COPinv, 9, 66, 73, 75, 204, 245, 246
- COPinv2, 9, 66, 74, 74, 247, 248
- densityCOP, 8, 76, 78, 79, 182, 219, 220, 224, 265, 266, 269, 290, 368, 372, 374
- densityCOPplot, 8, 76, 77, 78, 368, 370
- derCOP, 9–11, 81, 83, 84, 88, 107, 132, 133, 141, 149, 204, 211, 252, 253, 294, 318, 330, 355–357
- derCOP2, 9–11, 81, 82, 83, 89, 109, 129, 130, 149, 211, 251, 294, 318, 330, 355–357
- derCOPinv, 9, 82, 85, 110, 132, 211, 261, 300, 301, 327–330
- derCOPinv2, 9, 84, 89, 112, 129, 211, 264, 304, 305, 329, 330
- diagCOP, 7, 90, 93, 100, 137, 154, 319, 347
- diagCOPatf, 7, 28, 91, 92, 192, 194–196, 199
- diagCOPinv (diagCOPatf), 92
- duCOP, 7, 10–13, 53, 54, 68, 92, 94, 190, 192, 194, 195, 197, 199, 344

- EMPIRcop, [8](#), [16](#), [22](#), [23](#), [25](#), [37](#), [40](#), [41](#), [44](#), [97](#),
[102–104](#), [107–110](#), [113–115](#), [167](#),
[175](#), [206](#), [255](#), [257](#), [316](#), [317](#), [331](#),
[370](#), [376](#), [380](#), [382](#)
- EMPIRcopdf, [102](#), [104](#), [108–110](#), [113](#), [163](#)
- EMPIRgrid, [8](#), [103](#), [107–110](#), [112–115](#),
[125–128](#)
- EMPIRgrid_fast, [8](#), [98](#), [103](#), [104](#), [113](#), [115](#),
[375](#), [376](#), [381](#), [386](#)
- EMPIRgrid_lkup, [8](#), [104](#), [114](#), [115](#)
- EMPIRgridder, [107](#), [109](#), [110](#)
- EMPIRgridder2, [108](#), [109](#), [110](#), [113](#)
- EMPIRgridderinv, [110](#), [113](#), [117](#), [118](#),
[124–127](#)
- EMPIRgridderinv2, [9](#), [112](#), [113](#), [118](#), [124](#)
- EMPIRmed.regress, [116](#), [118](#), [124](#)
- EMPIRmed.regress2, [117](#), [117](#), [118](#), [124](#)
- EMPIRqua.regress, [117](#), [118](#), [118](#), [124](#)
- EMPIRqua.regress2, [117](#), [118](#), [124](#)
- EMPIRsim, [8](#), [125](#), [127](#), [128](#), [328](#)
- EMPIRsimv, [8](#), [125](#), [126](#), [127](#)
- EuvCOP, [10](#), [11](#), [129](#), [133](#), [251](#)
- EvuCOP, [10](#), [11](#), [130](#), [131](#), [132](#), [251–253](#)
- FGMcop, [6](#), [86](#), [134](#), [295](#)
- FGMicop (FGMcop), [134](#)
- footCOP, [7](#), [41](#), [100](#), [137](#), [155](#), [168](#), [186](#), [188](#),
[314](#), [357](#), [377](#)
- FRcop, [5](#), [6](#), [140](#), [202](#)
- FRECHETcop, [6](#), [47](#), [63](#), [143](#)
- gEvcop, [6](#), [145](#)
- GHcop, [6](#), [9](#), [12](#), [13](#), [67](#), [71](#), [147](#), [160](#), [170](#), [184](#),
[191](#), [196](#), [204](#), [208](#), [209](#), [229](#), [232](#),
[256](#), [271](#), [331](#), [334](#), [361](#), [362](#), [368](#)
- giniCOP, [7](#), [41](#), [71](#), [100](#), [139](#), [153](#), [168](#),
[173–175](#), [186](#), [188](#), [314](#), [322](#), [357](#),
[377](#)
- GLcop, [6](#), [152](#), [157](#), [170](#), [204](#), [342](#), [361](#), [362](#)
- GLEVcop (GLcop), [157](#)
- GLPMcop (GLcop), [157](#)
- glueCOP, [11](#), [47](#), [56](#), [59](#), [61](#), [63](#), [65](#), [66](#), [160](#),
[294](#), [296](#)
- gridCOP, [8](#), [162](#)
- HAIRPINcop, [6](#), [164](#)
- hoefCOP, [7](#), [41](#), [70](#), [99](#), [137](#), [139](#), [155](#), [165](#),
[186](#), [188](#), [255](#), [313](#), [314](#), [357](#), [377](#),
[392](#)
- HRcop, [6](#), [71](#), [152](#), [160](#), [169](#), [170](#), [204](#), [334](#),
[361](#), [362](#)
- isCOP.LTD, [10](#), [171](#), [175](#), [181](#)
- isCOP.permsym, [10](#), [31](#), [46](#), [58](#), [148](#), [166](#), [172](#),
[176–178](#), [186](#), [223](#), [259](#), [270](#), [375](#)
- isCOP.PQD, [10](#), [171](#), [172](#), [173](#), [180](#), [181](#), [374](#),
[377](#)
- isCOP.radsym, [10](#), [148](#), [172](#), [173](#), [176](#), [186](#)
- isCOP.RTI, [10](#), [172](#), [175](#), [179](#)
- isfuncCOP, [76](#), [181](#)
- J0copB5, [6](#), [182](#)
- J0copBB4 (GLcop), [157](#)
- joeskewCOP, [7](#), [8](#), [41](#), [100](#), [148](#), [155](#), [168](#), [184](#),
[314](#), [357](#), [365](#), [377](#)
- joint.curvesCOP, [9](#), [92](#), [93](#), [96](#), [189](#), [194](#),
[199](#), [246](#)
- joint.curvesCOP2, [9](#), [192](#), [193](#), [248](#)
- jointCOP, [7](#), [11](#), [92](#), [93](#), [96](#), [192](#), [194](#), [195](#)
- JOMAcop, [6](#), [200](#)
- kfuncCOP, [7](#), [12](#), [46](#), [203](#), [211](#), [212](#), [215](#), [217](#),
[356](#), [357](#), [373](#)
- kfuncCOP_Pd, [7](#), [216](#)
- kfuncCOPinv, [7](#), [204](#), [206](#), [211](#), [211](#), [212](#)
- kfuncCOPlmom, [7](#), [212](#)
- kfuncCOPlmom (kfuncCOPlmoms), [213](#)
- kfuncCOPlmoms, [7](#), [212](#), [213](#), [288](#)
- khoudraji1COP, [11](#)
- khoudraji1COP (composite1COP), [55](#)
- khoudraji2COP, [11](#)
- khoudraji2COP (composite2COP), [57](#)
- khoudrajiPCOP, [11](#)
- khoudrajiPCOP (composite1COP), [55](#)
- kmeasCOP, [7](#)
- kmeasCOP (kfuncCOP), [203](#)
- kullCOP, [10](#), [77](#), [218](#), [228](#), [269](#), [336](#), [342](#), [343](#),
[366](#), [368](#), [369](#), [372](#)
- kullCOPint (kullCOP), [218](#)
- lcomCOP, [7–9](#), [31](#), [35](#), [172](#), [177](#), [178](#), [222](#), [228](#),
[229](#), [232](#), [233](#), [271](#), [287](#), [323–326](#)
- lcomCOPpv, [10](#), [225](#), [226](#), [371](#)
- LCOMDIA_GH2cop, [9](#), [224](#), [229](#), [232](#), [233](#)
- LCOMDIA_GH3cop, [9](#), [224](#), [229](#), [232](#), [233](#)
- LCOMDIA_ManyCops, [9](#), [170](#), [224](#), [229](#), [232](#), [233](#)
- lcomoms2.ABcop2parameter, [238](#)
- lcomoms2.ABKGCop2parameter, [242](#)

- level.curvesCOP, 9, 73–75, 99, 100, 190, 192, 199, 245, 248, 249
- level.curvesCOP2, 9, 73–75, 193, 194, 246, 247, 249, 250
- level.setCOP, 9, 246, 248, 250
- level.setCOP2, 9, 248, 249, 249
- LMRuvCOP, 11, 250, 252, 253
- LMRvuCOP, 11, 251, 251
- LpCOP, 7
- LpCOP (hoefCOP), 165
- LpCOPpermsym, 255, 256, 259
- LpCOPpermsym (hoefCOP), 165
- LpCOPpradsym (hoefCOP), 165
- LzCOPpermsym, 7, 166, 168, 173, 254, 375
- M, 6, 11, 43, 46, 52, 53, 137, 140, 141, 143, 145, 149, 152, 154, 157, 160, 164, 165, 182–185, 196, 200, 202, 204, 207, 210, 245, 247, 260, 270, 273–276, 286–290, 294, 305, 307, 309, 310, 317, 318, 347, 359, 363, 364, 372, 373
- M_N5p12b, 272, 400
- med.regressCOP, 9, 13, 40, 132, 252, 261, 265, 300, 301
- med.regressCOP2, 9, 251, 264, 264, 304, 305
- m1eCOP, 8, 13, 76, 77, 135, 258, 265, 271, 335, 372
- MOcop, 5, 6, 9, 148, 232, 270
- N4212cop, 6, 207, 208, 274, 299, 300, 339
- N4220cop, 6, 275, 317
- NORMcop, 5, 6, 146, 202, 276, 359, 360
- nuskewCOP, 7, 8, 71, 151, 166, 191
- nuskewCOP (joeskewCOP), 184
- nustarCOP, 7, 8, 71, 166
- nustarCOP (joeskewCOP), 184
- ORDSUMcop, 6, 9, 11, 224, 233, 273, 278, 285, 377, 400
- ORDSUWcop, 6, 11, 278, 279, 284, 400
- P, 6, 8, 17, 19, 36, 43, 52, 53, 55, 58, 90, 134, 135, 139–141, 143, 145, 155, 157, 160, 165, 170, 173, 182–185, 198, 200, 202, 204, 206, 210, 216, 260, 261, 270, 271, 275, 276, 286, 287–289, 294, 299, 300, 305, 307, 309–311, 317, 353, 359, 372, 373, 377, 384, 391, 397
- PAcop, 6
- PAcop (PARETOcop), 287
- PARETOcop, 287
- PLACKETTcop, 207, 208, 289, 291–294, 342, 353, 369
- PLACKETTpar, 290, 291, 294
- PLACKETTsim, 290, 292, 293
- PLcop, 6, 177, 184, 287, 291, 292, 376
- PLcop (PLACKETTcop), 289
- PLpar, 290
- PLpar (PLACKETTpar), 291
- prod2COP, 11, 66, 294
- psepolar, 10, 297, 333, 334, 336, 340
- PSP, 6, 12, 44, 95, 132, 133, 177, 274, 299, 350, 369
- qua.regressCOP, 9, 261, 264, 300, 303
- qua.regressCOP.draw, 261, 264, 265, 301, 302
- qua.regressCOP2, 9, 264, 265, 303, 304
- RAYcop, 6, 305
- rCOP, 8
- rCOP (simCOP), 327
- ReineckeWell1266, 307
- ReineckeWells, 308
- RFcop, 6, 66, 309
- rhobevCOP, 148, 152, 311
- rhoCOP, 7, 9, 17, 19, 31–34, 40, 41, 100, 134, 138–140, 143, 145, 148, 155, 167, 168, 173–175, 177, 186–188, 200, 223, 262, 270, 273, 276, 287, 291, 292, 309, 311, 312, 313, 322, 342, 355, 357, 359, 363, 368, 374, 377, 399
- rmseCOP, 8, 16, 22, 23, 258, 316, 340, 343
- rN4220cop, 6, 275, 317
- sectionCOP, 9, 91, 318
- semicorCOP, 7, 8, 320
- simcomposite2COP (simcompositeCOP), 325
- simcomposite3COP, 61, 323, 326
- simcompositeCOP, 239, 240, 242, 244, 324, 325
- simCOP, 8, 9, 11, 61, 71, 72, 77, 79, 100, 125, 127, 141, 191, 240, 244, 275, 293, 318, 327, 329, 332, 369, 370, 372
- simCOPmicro, 8, 9, 66, 67, 127, 293, 328, 329, 344

simCOPv, 266
 simCOPv (simCOPmicro), 329
 spectralmeas, 10, 298, 333, 338, 340
 stabtaildepf, 10, 298, 336, 337
 statTn, 8, 16, 22, 316, 340, 372
 surCOP, 5, 7, 10, 12, 54, 67, 68, 94, 96, 166,
 176, 294, 330, 343, 345, 346, 353
 surfuncCOP, 7, 12, 43, 68, 344, 345, 353

 tailconCOP, 10, 347, 351, 354
 taildepCOP, 8, 10, 44, 45, 143, 148, 204, 207,
 271, 274, 289, 331, 348, 348, 354,
 370
 tailordCOP, 10, 348, 351, 353
 tauCOP, 7, 13, 17, 20, 40, 41, 52, 100, 139,
 140, 143, 155, 168, 173–175, 182,
 186, 188, 191, 200, 204, 209, 211,
 262, 263, 271, 273, 276, 292, 309,
 312–314, 322, 342, 355, 359, 363,
 368, 377, 391, 399
 Tcop, 5, 6, 276, 277, 358, 361, 362
 tEVcop, 6, 146, 152, 160, 170, 361
 TRIcop, 6, 363

 uvlmoms, 8, 35, 41, 100, 155, 166, 168, 185,
 225, 314, 357, 364, 377
 uvskew, 8, 10, 188
 uvskew (uvlmoms), 364

 vuongCOP, 10, 220, 228, 268, 328, 343, 349,
 366

 W, 6, 11, 43, 46, 52, 53, 140, 141, 143, 145,
 154, 165, 184, 185, 200, 202, 204,
 207, 210, 245, 247, 260, 261, 276,
 286, 287, 289, 290, 294, 347, 353,
 358, 363, 364, 372, 400
 W_N5p12a, 273, 279, 285, 399
 wolfCOP, 7, 8, 41, 100, 139, 155, 168, 174,
 186, 188, 314, 357, 374, 379–382,
 384, 387, 391, 392, 397
 wolfCOPsamc, 8, 375, 377, 379, 387
 wolfCOPtest, 8, 376, 377, 384, 392–394, 397
 wolfCOPtest_check, 391
 wolfCOPtest_data_smlsam, 384, 386, 387,
 397