Package 'NMOF'

January 20, 2025

Type Package

Title Numerical Methods and Optimization in Finance

Version 2.10-1

Date 2024-11-03

Maintainer Enrico Schumann <es@enricoschumann.net>

Depends R (>= 3.5)

Imports grDevices, graphics, parallel, stats, utils

Suggests MASS, PMwR, RUnit, Rglpk, datetimeutils, openxlsx, quadprog, readxl, tinytest, zoo

Description Functions, examples and data from the first and the second edition of ``Numerical Methods and Optimization in Finance" by M. Gilli, D. Maringer and E. Schumann (2019, ISBN:978-0128150658). The package provides implementations of optimisation heuristics (Differential Evolution, Genetic Algorithms, Particle Swarm Optimisation, Simulated Annealing and Threshold Accepting), and other optimisation tools, such as grid search and greedy search. There are also functions for the valuation of financial instruments such as bonds and options, for portfolio selection and functions that help with stochastic simulations.

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URL https://enricoschumann.net/NMOF.htm, https://gitlab.com/NMOF,

https://git.sr.ht/~enricoschumann/NMOF,

https://github.com/enricoschumann/NMOF

LazyLoad yes

LazyData yes

ByteCompile yes

Classification/JEL C61, C63

NeedsCompilation no

Author Enrico Schumann [aut, cre] (<https://orcid.org/0000-0001-7601-6576>)

Repository CRAN Date/Publication 2024-11-03 18:50:02 UTC

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NMOF-package

Numerical Methods and Optimization in Finance

Description

Functions, data and other R code from the book 'Numerical Methods and Optimization in Finance'. Comments/corrections/remarks/suggestions are very welcome (please contact the maintainer directly).

Details

The package contains implementations of several optimisation heuristics: Differential Evolution (DEopt), Genetic Algorithms (GAopt), (Stochastic) Local Search (LSopt), Particle Swarm (PSopt), Simuleated Annealing (SAopt) and Threshold Accepting (TAopt). The term heuristic is meant in the sense of general-purpose optimisation method.

Dependencies: The package is completely written in R. A number of packages are *suggested*, but they are not strictly required when using the **NMOF** package, and most of the package's functionality is available without them. Specifically, package **MASS** is needed to run the complete example for PSopt and also in one of the vignettes (PSIms). Package **parallel** is optional for functions bracketing, GAopt, gridSearch and restartOpt, and may become an option for other functions. Package **quadprog** is needed for a vignette (TAportfolio), some tests, and it may be used for computing mean-variance efficient portfolios. Package **Rglpk** is needed for function minCVaR. Package **readxl** is needed to process the raw data in function Shiller; package **datetimeutils** is used by French and Shiller. **PMwR** would be needed to run the examples of the backtesting examples in the NMOF book. Finally, packages **RUnit** and **tinytest** are needed to run the tests in subdirectory 'unitTests'.

Version numbering: package versions are numbered in the form major-minor-patch. The *patch* level is incremented with any published change in a version. *Minor* version numbers are incremented when a feature is added or an existing feature is substantially revised. (Such changes will be reported in the NEWS file.) The *major* version number will only be increased if there were a new edition of the book.

The source code of the **NMOF** package is also hosted at https://github.com/enricoschumann/ NMOF/. Updates to the package and new features are described at https://enricoschumann.net/ notes/NMOF/.

Optimisation:

There are functions for Differential Evolution (DEopt), Genetic Algorithms (GAopt), (Stochastic) Local Search (LSopt), Simuleated Annealing (SAopt), Particle Swarm (SAopt), and Threshold Accepting (TAopt). The function restartOpt helps with running restarts of these methods; also available are functions for grid search (gridSearch) and greedy search (greedySearch).

Pricing Financial Instruments:

For options: See vanillaOptionEuropean, vanillaOptionAmerican, putCallParity. For pricing methods that use the characteristic function, see callCF.

For bonds and bond futures: See vanillaBond, bundFuture and xtContractValue.

Simulation:

See resampleC and mc.

Data:

See bundData, fundData and optionData.

Author(s)

Enrico Schumann

Maintainer: Enrico Schumann <es@enricoschumann.net>

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

Examples

```
## Not run:
library("NMOF")
```

overview
packageDescription("NMOF")
help(package = "NMOF")

```
## code from book
showExample("equations.R", edition = 1)
showExample("Heur")
```

```
## show NEWS file
news(Version >= "2.0-0", package = "NMOF")
```

```
## vignettes
vignette(package = "NMOF")
nss <- vignette("DEnss", package = "NMOF")
print(nss)
edit(nss)</pre>
```

```
## _book_ websites
browseURL("https://nmof.net")
browseURL("https://enricoschumann.net/NMOF/")
## _package_ websites
browseURL("https://enricoschumann.net/R/packages/NMOF/")
browseURL("https://cran.r-project.org/package=NMOF")
browseURL("https://git.sr.ht/~enricoschumann/NMOF")
browseURL("https://github.com/enricoschumann/NMOF")
## unit tests
file.show(system.file("unitTests/test_results.txt", package = "NMOF"))
## End(Not run)
test.rep <- readLines(system.file("unitTests/test_results.txt",</pre>
                                   package = "NMOF"))
nt <- gsub(".*\\(([0-9]+) checks?\\).*", "\\1",</pre>
           test.rep[grep("\\(\\d+ checks?\\)", test.rep)])
message("Number of unit tests: ", sum(as.numeric(nt)))
```

approxBondReturn Approximate Total Return of Bond

Description

Approximate the total return of a bond by its current yield, duration and convexity.

Usage

```
approxBondReturn(yield, tm, n = 2, scale = 1/250, pad = NULL)
```

Arguments

yield	a numeric vector
tm	a numeric vector: time-to-maturity
n	number of coupon payments per period
scale	how to scale yield; see Details
pad	how to pad the first observation: NULL (default) means to drop it; useful alternatives are NA or ${\tt 0}$

Details

The function approximates the total return of a bond investor, based on changes in yield. The computation is based on a Taylor-series expansion. See the references, in particular concerning the shortcomings of the approximation:

- 1. approximation is based on par yield
- 2. it relies on yield alone, so does not take into account defaults; so for indices, the approximation should only used for issuers without defaults

Value

a numeric vector, with attributes duration and convexity

Note

Package treasuryTR implements the method as well.

Author(s)

Enrico Schumann

References

Swinkels, L. (2019). Treasury Bond Return Data Starting in 1962. Data. 4 (3).

Tuckman, B. and Serrat, A. (2012). Fixed Income Securities – Tools for Today's Markets. 3rd edition. Wiley.

Examples

```
yield0 <- 0.05
tm <- 20
cf <- c(rep(5, tm-1), 105)
duration(cf, 1:tm, yield0)
approxBondReturn(yield = c(yield0, 0.05), tm = tm, n = 1)
## ==> no price change, current yield is earned
approxBondReturn(yield = c(yield0, 0.04), tm = tm, n = 1)
## ==> current yield + price changed is earned
```

bracketing Zero-Bracketing

Description

Bracket the zeros (roots) of a univariate function

Usage

```
bracketing(fun, interval, ...,
    lower = min(interval), upper = max(interval),
    n = 20L,
    method = c("loop", "vectorised", "multicore", "snow"),
    mc.control = list(), cl = NULL)
```

bracketing

Arguments

fun	a univariate function; it will be called as fun(x,) with x being a numeric vector
interval	a numeric vector, containing the end-points of the interval to be searched
	further arguments passed to fun
lower	lower end-point. Ignored if interval is specified.
upper	upper end-point. Ignored if interval is specified.
n	the number of function evaluations. Must be at least 2 (in which case fun is evaluated only at the end-points); defaults to 20.
method	can be loop (the default), vectorised, multicore or snow. See Details.
mc.control	a list containing settings that will be passed to mclapply if method is multicore. Must be a list of named elements. See the documentation of mclapply in pack- age parallel .
cl	default is NULL. If method is snow, this must be a cluster object or an integer (the number of cores to be used). See the documentation of packages parallel and snow .

Details

bracketing evaluates fun at equal-spaced values of x between (and including) lower and upper. If the sign of fun changes between two consecutive x-values, bracketing reports these two x-values as containing ('bracketing') a root. There is no guarantee that there is only one root within a reported interval. bracketing will not narrow the chosen intervals.

The argument method determines how fun is evaluated. Default is loop. If method is "vectorised", fun must be written such that it can be evaluated for a vector x (see Examples). If method is multicore, function mclapply from package **parallel** is used. Further settings for mclapply can be passed through the list mc.control. If multicore is chosen but the functionality is not available (eg, currently on Windows), then method will be set to loop and a warning is issued. If method is snow, function clusterApply from package **parallel** is used. In this case, the argument cl must either be a cluster object (see the documentation of clusterApply) or an integer. If an integer, a cluster will be set up via makeCluster(c(rep("localhost", cl)), type = "SOCK"), and stopCluster is called when the function is exited. If snow is chosen but the package is not available or cl is not specified, then method will be set to loop and a warning is issued. In case that cl is a cluster object, stopCluster will not be called automatically.

Value

A numeric matrix with two columns, named *lower* and *upper*. Each row contains one interval that contains at least one root. If no roots were found, the matrix has zero rows.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

See Also

uniroot (in package stats)

Examples

```
## Gilli/Maringer/Schumann (2011), p. 290
testFun <- function(x)
    cos(1/x^2)
bracketing(testFun, interval = c(0.3, 0.9), n = 26L)
bracketing(testFun, interval = c(0.3, 0.9), n = 26L, method = "vectorised")</pre>
```

bundData

German Government Bond Data

Description

A sample of data on 44 German government bonds. Contains ISIN, coupon, maturity and dirty price as of 2010-05-31.

Usage

bundData

Format

bundData is a list with three components: cfList, tmList and bM. cfList is list of 44 numeric vectors (the cash flows). tmList is a list of 44 character vectors (the payment dates) formatted as YYYY-MM-DD. bM is a numeric vector with 44 elements (the dirty prices of the bonds).

Details

All prices are as of 31 May 2010. See chapter 14 in Gilli et al. (2011).

Source

The data was obtained from https://www.deutsche-finanzagentur.de/en/. The data is also freely available from the website of the Bundesbank https://www.bundesbank.de/en/.

bundFuture

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

Examples

```
bundFuture
```

Theoretical Valuation of Euro Bund Future

Description

Compute theoretical prices of bund future.

Usage

Arguments

clean	numeric: clean prices of CTD
future	numeric: price of future
coupon	numeric
trade.date	Date or character in format YYYY-MM-DD
expiry.date	Date or character in format YYYY-MM-DD
last.coupon.dat	te
	Date or character in format YYYY-MM-DD
r	numeric: 0.01
cf	numeric: conversion factor of CTD

Details

bundFuture computes the theoretical prices of the Bund Future, given the prices of the cheapestto-deliver eligible government bond.

bundFutureImpliedRate computes the implied refinancing rate.

Value

numeric

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Examples

```
## Bund-Future with expiry Sep 2017
## CTD: DE0001102408 -- 0%, 15 Aug 2026
##
## On 21 August 2017, the CTD traded (clean) at 97.769
## the FGBL Sep 2017 closed at 164.44.
                                             ## DE0001102408
bundFuture(clean = 97.769,
           coupon = 0,
           trade.date = "2017-8-21",
           expiry.date = "2017-09-07",
                                             ## Bund expiry
           last.coupon.date = "2017-08-15", ## last co
           r = -0.0037,
           cf = 0.594455)
                            ## conversion factor (from Eurex website)
bundFutureImpliedRate(future = 164.44,
                      clean = 97.769,
                      coupon = 0,
                      trade.date = "2017-8-21",
                      expiry.date = "2017-09-07",
                      last.coupon.date = "2017-08-15",
                      cf = 0.594455)
```

callCF

Description

Price a European plain-vanilla call with the characteric function.

Usage

Arguments

cf	characteristic function
S	spot
Х	strike
tau	time to maturity
r	the interest rate
q	the dividend rate
	arguments passed to the characteristic function
implVol	logical: compute implied vol?
uniroot.contro	\mathbf{p}
	A list. If there are elements named interval, tol or maxiter, these are passed to uniroot. Any other elements of the list are ignored.
uniroot.info	logical; default is FALSE. If TRUE, the function will return the information re- turned by uniroot. See paragraph Value below.
om	a (usually complex) argument
vØ	a numeric vector of length one
vT	a numeric vector of length one
v	a numeric vector of length one
rho	a numeric vector of length one
k	a numeric vector of length one
sigma	a numeric vector of length one
lambda	a numeric vector of length one
muJ	a numeric vector of length one
vJ	a numeric vector of length one
nu	a numeric vector of length one
theta	a numeric vector of length one

Details

The function computes the value of a plain vanilla European call under different models, using the representation of Bakshi/Madan. Put values can be computed through put–call parity (see putCallParity).

If implVol is TRUE, the function will compute the implied volatility necessary to obtain the same value under Black–Scholes–Merton. The implied volatility is computed with uniroot from the **stats** package. The default search interval is c(0.00001, 2); it can be changed through uniroot.control.

The function uses variances as inputs (not volatilities).

The function is not vectorised (but see the NMOF Manual for examples of how to efficiently price more than one option at once).

Value

Returns the value of the call (numeric) under the respective model or, if implVol is TRUE, a list of the value and the implied volatility. (If, in addition, uniroot.info is TRUE, the information provided by uniroot is also returned.)

Note

If implVol is TRUE, the function will return a list with elements named value and impliedVol. Prior to version 0.26-3, the first element was named callPrice.

Author(s)

Enrico Schumann

References

Bates, David S. (1996) Jumps and Stochastic Volatility: Exchange Rate Processes Implicit in Deutsche Mark Options. *Review of Financial Studies* **9** (1), 69–107.

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Heston, S.L. (1993) A Closed-Form Solution for Options with Stochastic Volatility with Applications to Bonds and Currency options. *Review of Financial Studies* **6** (2), 327–343.

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

callHestoncf

Examples

```
S <- 100; X <- 100; tau <- 1
r <- 0.02; q <- 0.08
v0 <- 0.2^2 ## variance, not volatility
vT <- 0.2^2 ## variance, not volatility
v <- vT</pre>
```

callHestoncf

```
rho <- -0.3; k <- .2
sigma <- 0.3
## jump parameters (Merton and Bates)
lambda <- 0.1
muJ <- -0.2
vJ <- 0.1^2
## get Heston price and BSM implied volatility
callHestoncf(S, X, tau, r, q, v0, vT, rho, k, sigma, implVol = FALSE)
callCF(cf = cfHeston, S=S, X=X, tau=tau, r=r, q = q,
       v0 = v0, vT = vT, rho = rho, k = k, sigma = sigma, implVol = FALSE)
## Black-Scholes-Merton
callCF(cf = cfBSM, S=S, X=X, tau = tau, r = r, q = q,
       v = v, implVol = TRUE)
## Bates
callCF(cf = cfBates, S = S, X = X, tau = tau, r = r, q = q,
       v0 = v0, vT = vT, rho = rho, k = k, sigma = sigma,
       lambda = lambda, muJ = muJ, vJ = vJ, implVol = FALSE)
## Merton
callCF(cf = cfMerton, S = S, X = X, tau = tau, r = r, q = q,
       v = v, lambda = lambda, muJ = muJ, vJ = vJ, implVol = FALSE)
## variance gamma
nu <- 0.1; theta <- -0.1; sigma <- 0.15
callCF(cf = cfVG, S = S, X = X, tau = tau, r = r, q = q,
       nu = nu, theta = theta, sigma = sigma, implVol = FALSE)
```

callHestoncf

Price of a European Call under the Heston Model

Description

Computes the price of a European Call under the Heston model (and the equivalent Black–Scholes– Merton volatility)

Usage

Arguments

S	current stock price
Х	strike price

tau	time to maturity
r	risk-free rate
q	dividend rate
v0	current variance
vT	long-run variance (theta in Heston's paper)
rho	correlation between spot and variance
k	speed of mean-reversion (kappa in Heston's paper)
sigma	volatility of variance. A value smaller than 0.01 is replaced with 0.01.
implVol	compute equivalent Black-Scholes-Merton volatility? Default is FALSE.
	named arguments, passed to integrate
uniroot.contro	1
	A list. If there are elements named interval, tol or maxiter, these are passed to uniroot. Other elements of the list are ignored.
uniroot.info	logical; default is FALSE. If TRUE, the function will return the information re- turned by uniroot. See section Value below.

Details

The function computes the value of a plain vanilla European call under the Heston model. Put values can be computed through put–call-parity.

If implVol is TRUE, the function will compute the implied volatility necessary to obtain the same price under Black–Scholes–Merton. The implied volatility is computed with uniroot from the stats package (the default search interval is c(0.00001, 2); it can be changed through uniroot.control).

Note that the function takes variances as inputs (not volatilities).

Value

Returns the value of the call (numeric) under the Heston model or, if implVol is TRUE, a list of the value and the implied volatility. If uniroot.info is TRUE, then instead of only the computed volatility, the complete output of uniroot is included in the result.

Note

If implVol is TRUE, the function will return a list with elements named value and impliedVol. Prior to version 0.26-3, the first element was named callPrice.

Author(s)

Enrico Schumann

callMerton

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Heston, S.L. (1993) A Closed-Form Solution for Options with Stochastic Volatility with Applications to Bonds and Currency options. *Review of Financial Studies* **6**(2), 327–343.

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

callCF, EuropeanCall

Examples

```
S <- 100; X <- 100; tau <- 1; r <- 0.02; q <- 0.01
v0 <- 0.2<sup>2</sup> ## variance, not volatility
vT <- 0.2<sup>2</sup> ## variance, not volatility
rho <- -0.7; k <- 0.2; sigma <- 0.5
## get Heston price and BSM implied volatility
result <- callHestoncf(S = S, X = X, tau = tau, r = r, q = q,</pre>
                        v0 = v0, vT = vT, rho = rho, k = k,
                        sigma = sigma, implVol = TRUE)
## Heston price
result[[1L]]
## price BSM with implied volatility
vol <- result[[2L]]</pre>
d1 <- (log(S/X) + (r - q + vol^2 / 2)*tau) / (vol*sqrt(tau))</pre>
d2 <- d1 - vol*sqrt(tau)
callBSM <- S * exp(-q * tau) * pnorm(d1) -</pre>
           X * exp(-r * tau) * pnorm(d2)
callBSM ## should be (about) the same as result[[1L]]
```

```
callMerton
```

Price of a European Call under Merton's Jump-Diffusion Model

Description

Computes the price of a European Call under Merton's jump-diffusion model (and the equivalent Black-Scholes-Merton volatility)

Usage

```
callMerton(S, X, tau, r, q, v, lambda, muJ, vJ, N, implVol = FALSE)
```

callMerton

Arguments

S	current stock price
Х	strike price
tau	time to maturity
r	risk-free rate
q	dividend rate
v	variance
lambda	jump intensity
muJ	mean jump-size
vJ	variance of log jump-size
Ν	The number of jumps. See Details.
implVol	compute equivalent Black-Scholes-Merton volatility? Default is FALSE.

Details

The function computes the value of a plain-vanilla European call under Merton's jump-diffusion model. Put values can be computed through put-call-parity (see putCallParity). If implVol is TRUE, the function also computes the implied volatility necessary to obtain the same price under Black-Scholes-Merton. The implied volatility is computed with uniroot from the stats package.

Note that the function takes variances as inputs (not volatilities).

The number of jumps N typically can be set 10 or 20. (Just try to increase N and see how the results change.)

Value

Returns the value of the call (numeric) or, if implVol is TRUE, a list of the value and the implied volatility.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Merton, R.C. (1976) Option Pricing when Underlying Stock Returns are Discontinuous. *Journal of Financial Economics* **3**(1–2), 125–144.

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

See Also

callCF, EuropeanCall

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colSubset

Examples

```
S <- 100; X <- 100; tau <- 1
r <- 0.0075; q <- 0.00
v <- 0.2^2
lambda <- 1; muJ <- -0.2; vJ <- 0.6^2
N <- 20
## jumps can make a difference
callMerton(S, X, tau, r, q, v, lambda, muJ, vJ, N, implVol = TRUE)
callCF(cf = cfMerton, S = S, X = X, tau = tau, r = r, q = q,
       v = v, lambda = lambda, muJ = muJ, vJ = vJ, implVol = TRUE)
vanillaOptionEuropean(S,X,tau,r,q,v, greeks = FALSE)
lambda <- 0 ## no jumps
callMerton(S, X, tau, r, q, v, lambda, muJ, vJ, N, implVol = FALSE)
vanillaOptionEuropean(S,X,tau,r,q,v, greeks = FALSE)
lambda <- 1; muJ <- 0; vJ <- 0.0^2 ## no jumps, either
callMerton(S, X, tau, r, q, v, lambda, muJ, vJ, N, implVol = FALSE)
vanillaOptionEuropean(S,X,tau,r,q,v, greeks = FALSE)
```

colSubset

Full-rank Column Subset

Description

Select a full-rank subset of columns of a matrix.

Usage

```
colSubset(x)
```

Arguments

х

a numeric matrix

Details

Uses qr.

Value

A list:

columns	indices of columns
multiplier	a matrix

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

repairMatrix

Examples

```
nc <- 3
           ## columns
nr <- 10 ## rows
M <- array(rnorm(nr * nc), dim = c(nr, nc))</pre>
C \leq array(0.5, dim = c(nc, nc))
diag(C) <- 1
M <- M %*% chol(C)
M <- M[ ,c(1,1,1,2,3)]
М
(tmp <- colSubset(M))</pre>
C <- cor(M[ ,tmp$columns])</pre>
nc <- ncol(C)</pre>
nr <- 100
X <- array(rnorm(nr*nc), dim = c(nr, nc))</pre>
X <- X %*% chol(C)
X <- X %*% tmp$multiplier
head(X)
cor(X)
```

CPPI

Constant-Proportion Portfolio Insurance

Description

Simulate constant-proportion portfolio insurance (CPPI) for a given price path.

Usage

```
CPPI(S, multiplier, floor, r, tau = 1, gap = 1)
```

CPPI

Arguments

S	numeric: price path of risky asset
multiplier	numeric
floor	numeric: a percentage, should be smaller than 1
r	numeric: interest rate (per time period tau)
tau	numeric: time periods
gap	numeric: how often to rebalance. 1 means every timestep, 2 means every second timestep, and so on.

Details

Based on Dietmar Maringer's MATLAB code (function CPPIgap, Listing 9.1). See Gilli, Maringer and Schumann, 2011, chapter 9.

Value

A list:	
V	normalised value (always starts at 1)
С	cushion
В	bond investment
F	floor
E	exposure
Ν	units of risky asset
S	price path

Author(s)

Original MATLAB code: Dietmar Maringer. R implementation: Enrico Schumann.

References

Chapter 9 of Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). $\label{eq:MOFManual} https://enricoschumann.net/NMOF.htm#NMOFmanual$

Examples

```
par(mfrow = c(3,1), mar = c(3,3,1,1))
plot(0:(length(S)-1), S, type = "s", main = "stock price")
plot(0:(length(S)-1), sol$V, type = "s", main = "value")
plot(0:(length(S)-1), 100*sol$E/sol$V, type = "s",
    main = "% invested in risky asset")
## rebalancing every 5th day
sol <- CPPI(S, multiplier = 5, floor = 0.9, r = 0.01,
        tau = tau, gap = 5)
par(mfrow = c(3,1), mar = c(3,3,1,1))
plot(0:(length(S)-1), S, type = "s", main = "stock price")
plot(0:(length(S)-1), sol$V, type = "s", main = "value")
plot(0:(length(S)-1), 100*sol$E/sol$V, type = "s",
        main = "% invested in risky asset")</pre>
```

```
DEopt
```

Optimisation with Differential Evolution

Description

The function implements the standard Differential Evolution algorithm.

Usage

DEopt(OF, algo = list(), ...)

Arguments

OF	The objective function, to be minimised. See Details.
algo	A list with the settings for algorithm. See Details and Examples.
	Other pieces of data required to evaluate the objective function. See Details and Examples.

Details

The function implements the standard Differential Evolution (no jittering or other features). Differential Evolution (DE) is a population-based optimisation heuristic proposed by Storn and Price (1997). DE evolves several solutions (collected in the 'population') over a number of iterations ('generations'). In a given generation, new solutions are created and evaluated; better solutions replace inferior ones in the population. Finally, the best solution of the population is returned. See the references for more details on the mechanisms.

To allow for constraints, the evaluation works as follows: after a new solution is created, it is (i) repaired, (ii) evaluated through the objective function, (iii) penalised. Step (ii) is done by a call to OF; steps (i) and (iii) by calls to algo\$repair and algo\$pen. Step (i) and (iii) are optional, so the respective functions default to NULL. A penalty is a positive number added to the 'clean' objective function value, so it can also be directly written in the OF. Writing a separate penalty function is often clearer; it can be more efficient if either only the objective function or only the penalty function

DEopt

can be vectorised. (Constraints can also be added without these mechanisms. Solutions that violate constraints can, for instance, be mapped to feasible solutions, but without actually changing them. See Maringer and Oyewumi, 2007, for an example.)

Conceptually, DE consists of two loops: one loop across the generations and, in any given generation, one loop across the solutions. DEopt indeed uses, as the default, two loops. But it does not matter in what order the solutions are evaluated (or repaired or penalised), so the second loop can be vectorised. This is controlled by the variables algo\$loopOF, algo\$loopRepair and algo\$loopPen, which all default to TRUE. Examples are given in the vignettes and in the book. The respective algo\$loopFun must then be set to FALSE.

All objects that are passed through . . . will be passed to the objective function, to the repair function and to the penalty function.

The list algo collects the the settings for the algorithm. Strictly necessary are only min and max (to initialise the population). Here are all possible arguments:

CR probability for crossover. Defaults to 0.9. Using default settings may not be a good idea.

- F The step size. Typically a numeric vector of length one; default is 0.5. Using default settings may not be a good idea. (F can also be a vector with different values for each decision variable.)
- nP population size. Defaults to 50. Using default settings may not be a good idea.
- nG number of generations. Defaults to 300. Using default settings may not be a good idea.
- min, max vectors of minimum and maximum parameter values. The vectors min and max are used to determine the dimension of the problem and to randomly initialise the population. Per default, they are no constraints: a solution may well be outside these limits. Only if algo\$minmaxConstr is TRUE will the algorithm repair solutions outside the min and max range.

minmaxConstr if TRUE, algo\$min and algo\$max are considered constraints. Default is FALSE.

- pen a penalty function. Default is NULL (no penalty).
- initP optional: the initial population. A matrix of size length(algo\$min) times algo\$nP, or a function that creates such a matrix. If a function, it should take no arguments.
- repair a repair function. Default is NULL (no repairing).
- loopOF logical. Should the OF be evaluated through a loop? Defaults to TRUE.
- loopPen logical. Should the penalty function (if specified) be evaluated through a loop? Defaults to TRUE.
- loopRepair logical. Should the repair function (if specified) be evaluated through a loop? Defaults to TRUE.
- printDetail If TRUE (the default), information is printed. If an integer i greater then one, information is printed at very ith generation.
- printBar If TRUE (the default), a txtProgressBar is printed.
- storeF if TRUE (the default), the objective function values for every solution in every generation are stored and returned as matrix Fmat.
- storeSolutions default is FALSE. If TRUE, the solutions (ie, decision variables) in every generation
 are stored and returned as a list P in list xlist (see Value section below). To check, for
 instance, the solutions at the end of the ith generation, retrieve xlist[[c(1L, i)]]. This will
 be a matrix of size length(algo\$min) times algo\$nP. (To be consistent with other functions,
 xlist is itself a list. In the case of DEopt, it contains just one element.)

- classify Logical; default is FALSE. If TRUE, the result will have a class attribute TAopt attached. This feature is **experimental**: the supported methods may change without warning.
- drop If FALSE (the default), the dimension is not dropped from a single solution when it is passed to a function. (That is, the function will receive a single-column matrix.)

Value

A list:

xbest	the solution (the best member of the population), which is a numeric vector
OFvalue	objective function value of best solution
рорҒ	a vector. The objective function values in the final population.
Fmat	if algo\$storeF is TRUE, a matrix of size algo\$nG times algo\$nP containing the objective function values of all solutions over the generations; else NA.
xlist	if algo\$storeSolutions is TRUE, a list that contains a list P of matrices and a matrix initP (the initial solution); else NA.
initial.state	the value of .Random. seed when the function was called.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Maringer, D. and Oyewumi, O. (2007). Index Tracking with Constrained Portfolios. *Intelligent Systems in Accounting, Finance and Management*, **15**(1), pp. 57–71.

Schumann, E. (2012) Remarks on 'A comparison of some heuristic optimization methods'. https://enricoschumann.net/R/remarks.htm

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Storn, R., and Price, K. (1997) Differential Evolution – a Simple and Efficient Heuristic for Global Optimization over Continuous Spaces. *Journal of Global Optimization*, **11**(4), pp. 341–359.

See Also

GAopt, PSopt

Examples

DEopt

```
F = 0.6,
                                 ### step size
             CR = 0.9,
                                 ### prob of crossover
            min = c(-10, -10), ### range for initial population
            max = c(10, 10))
sol <- DEopt(OF = OF, algo = algo)</pre>
## correct answer: -3.30686864747523
format(sol$OFvalue, digits = 12)
## check convergence of population
sd(sol$popF)
ts.plot(sol$Fmat, xlab = "generations", ylab = "OF")
## Example 2: vectorising the evaluation of the population
OF <- tfRosenbrock
                       ### see ?testFunctions
size <- 3L
                        ### define dimension
x <- rep.int(1, size) ### the known solution ...</pre>
0F(x)
                       ### ... should give zero
algo <- list(printBar = FALSE,</pre>
                   nP = 30L,
                    nG = 300L,
                    F = 0.6,
                   CR = 0.9,
                   min = rep(-100, size),
                   max = rep( 100, size))
## run DEopt
(t1 <- system.time(sol <- DEopt(OF = OF, algo = algo)))</pre>
sol$xbest
sol$OFvalue ### should be zero (with luck)
## a vectorised Rosenbrock function: works only with a *matrix* x
OF2 <- function(x) {
    n <- dim(x)[1L]</pre>
    xi <- x[seq_len(n - 1L), ]</pre>
    colSums(100 * (x[2L:n, ] - xi * xi)<sup>2</sup> + (1 - xi)<sup>2</sup>)
}
## random solutions (every column of 'x' is one solution)
x <- matrix(rnorm(size * algo$nP), size, algo$nP)</pre>
all.equal(OF2(x)[1:3],
          c(OF(x[ ,1L]), OF(x[ ,2L]), OF(x[ ,3L])))
## run DEopt and compare computing time
algo$loopOF <- FALSE
(t2 <- system.time(sol2 <- DEopt(OF = OF2, algo = algo)))</pre>
sol2$xbest
sol2$0Fvalue
                   ### should be zero (with luck)
t1[[3L]]/t2[[3L]] ### speedup
```

divRatio

Description

Compute the diversification ratio of a portfolio.

Usage

divRatio(w, var)

Arguments

W	numeric: a vector of weights
var	numeric matrix: the variance-covariance matrix

Details

The function provides an efficient implementation of the diversification ratio, suitable for optimisation.

Value

a numeric vector of length one

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Yves Choueifaty and Yves Coignard (2008) Toward Maximum Diversification. *Journal of Portfolio Management* **35**(1), 40–51.

See Also

pm, drawdown

Examples

na <- 10 ## number of assets
rho <- 0.5 ## correlation
v_min <- 0.2 ## minimum vol
v_max <- 0.4 ## maximum vol
set up a covariance matrix S</pre>

drawdown

```
C <- array(rho, dim = c(na,na))
diag(C) <- 1
vols <- seq(v_min, v_max, length.out = na)
S <- outer(vols, vols) * C
w <- rep(1/na, na) ## weights
divRatio(w, S)</pre>
```

drawdown

Drawdown

Description

Compute the drawdown of a time series.

Usage

drawdown(v, relative = TRUE, summary = TRUE)

Arguments

v	a price series (a numeric vector)
relative	if TRUE, maximum drawdown is chosen according to percentage losses; else in units of $\boldsymbol{\nu}$
summary	if TRUE, provide maximum drawdown and time when it occured; else return drawdown vector

Details

The drawdown at position t of a time series v is the difference between the highest peak that was reached before t and the current value. If the current value represents a new high, the drawdown is zero.

Value

If summary is FALSE, a vector of the same length as v. If summary is TRUE, a list

maximum	maximum drawdown
high	the max of v
high.position	position of high
low	the min of v
low.position	position of low

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

See Also

drawdowns

Examples

```
v <- cumprod(1 + rnorm(20) * 0.02)
drawdown(v)</pre>
```

EuropeanCall Computing Prices of European Calls with a Binomial Tree

Description

Computes the fair value of a European Call with the binomial tree of Cox, Ross and Rubinstein.

Usage

EuropeanCall(S0, X, r, tau, sigma, M = 101)
EuropeanCallBE(S0, X, r, tau, sigma, M = 101)

Arguments

S0	current stock price
Х	strike price
r	risk-free rate
tau	time to maturity
sigma	volatility
М	number of time steps

Details

Prices a European Call with the tree approach of Cox, Ross, Rubinstein.

The algorithm in EuropeanCallBE does not construct and traverse a tree, but computes the terminal prices via a binomial expansion (see Higham, 2002, and Chapter 5 in Gilli/Maringer/Schumann, 2011).

Value

Returns the value of the call (numeric).

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French

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

M. Gilli and Schumann, E. (2009) Implementing Binomial Trees. COMISEF Working Paper Series No. 008. https://enricoschumann.net/COMISEF/wps008.pdf

Higham, D. (2002) Nine Ways to Implement the Binomial Method for Option Valuation in MAT-LAB. *SIAM Review*, **44**(4), pp. 661–677. doi:10.1137/S0036144501393266.

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

See Also

callHestoncf

Examples

```
French
```

Download Datasets from Kenneth French's Data Library

Description

Download datasets from Kenneth French's Data Library.

Usage

Arguments

dest.dir	string: a path to a directory	
dataset	a character string: the CSV file name. Also supported are the keywords 'market' and 'rf'.	
weighting	a character string: "equal" or "value"	
frequency	a character string: daily, monthly or annual. Whether it is used or ignored depends on the particular dataset.	
price.series	logical: convert the returns series into prices series?	
na.rm	logical: remove missing values in the calculation of price series?	
adjust.frequency		
	logical: if TRUE, frequency is switched to '"daily"' when the word '"daily"' appears in the dataset's name	
return.class	a string: 'data.frame' (the default) or 'zoo'	

Details

The function downloads data provided by Kenneth French at https://mba.tuck.dartmouth.edu/ pages/faculty/ken.french/data_library.html. The download file gets a date prefix (current date in format YYYYMMDD) and is stored in directory dest.dir. Before any download is attempted, the function checks whether a file with today's prefix exist in dest.dir; if yes, the file is used.

In the original data files, missing values are coded as -99 or similar. These numeric values are replaced by NA.

Calling the function without any arguments will print the names of the supported datasets (and return them insivibly).

For dataset 'market', the function downloads the three-factor dataset, and adds excess return and risk-free rate. For dataset 'rf', only the risk-free rate is returned.

Value

A data.frame, with contents depending on the particular dataset, or an object as specified by return.class.

If the download fails, the function evaluates to NULL, typically with warnings.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

fundData

See Also

Shiller

Examples

list all supported files
French()

```
## Not run:
archive.dir <- "~/Downloads/French"
if (!dir.exists(archive.dir))
    dir.create(archive.dir)
French(archive.dir, "F-F_Research_Data_Factors_CSV.zip")
```

End(Not run)

fundData

Mutual Fund Returns

Description

A matrix of 500 rows (return scenarios) and 200 columns (mutual funds). The elements in the matrix are weekly returns.

Usage

fundData

Format

A plain numeric matrix.

Details

The scenarios were created with a bootstrapping technique. The data set is only meant to provide example data on which to test algorithms.

Source

Schumann, E. (2010) *Essays on Practical Financial Optimisation*, (chapter 4), PhD thesis, University of Geneva.

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

Examples

apply(fundData, 2, summary)

```
GAopt
```

Optimisation with a Genetic Algorithm

Description

A simple Genetic Algorithm for minimising a function.

Usage

GAopt (OF, algo = list(), ...)

Arguments

OF	The objective function, to be minimised. See Details.
algo	A list with the settings for algorithm. See Details and Examples.
	Other pieces of data required to evaluate the objective function. See Details and Examples.

Details

The function implements a simple Genetic Algorithm (GA). A GA evolves a collection of solutions (the so-called population), all of which are coded as vectors containing only zeros and ones. (In GAopt, solutions are of mode logical.) The algorithm starts with randomly-chosen or usersupplied population and aims to iteratively improve this population by mixing solutions and by switching single bits in solutions, both at random. In each iteration, such randomly-changed solutions are compared with the original population and better solutions replace inferior ones. In GAopt, the population size is kept constant.

GA language: iterations are called generations; new solutions are called offspring or children (and the existing solutions, from which the children are created, are parents); the objective function is called a fitness function; mixing solutions is a crossover; and randomly changing solutions is called mutation. The choice which solutions remain in the population and which ones are discarded is

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GAopt

called selection. In GAopt, selection is pairwise: a given child is compared with a given parent; the better of the two is kept. In this way, the best solution is automatically retained in the population.

To allow for constraints, the evaluation works as follows: after new solutions are created, they are (i) repaired, (ii) evaluated through the objective function, (iii) penalised. Step (ii) is done by a call to OF; steps (i) and (iii) by calls to algo\$repair and algo\$pen. Step (i) and (iii) are optional, so the respective functions default to NULL. A penalty can also be directly written in the OF, since it amounts to a positive number added to the 'clean' objective function value; but a separate function is often clearer. A separate penalty function is advantagous if either only the objective function or only the penalty function can be vectorised.

Conceptually a GA consists of two loops: one loop across the generations and, in any given generation, one loop across the solutions. This is the default, controlled by the variables algo\$loopOF, algo\$loopRepair and algo\$loopPen, which all default to TRUE. But it does not matter in what order the solutions are evaluated (or repaired or penalised), so the second loop can be vectorised. The respective algo\$loopFun must then be set to FALSE. (See also the examples for DEopt and PSopt.)

The evaluation of the objective function in a given generation can even be distributed. For this, an argument algo\$methodOF needs to be set; see below for details (and Schumann, 2011, for examples).

All objects that are passed through . . . will be passed to the objective function, to the repair function and to the penalty function.

The list algo contains the following items:

- nB number of bits per solution. Must be specified.
- nP population size. Defaults to 50. Using default settings may not be a good idea.
- nG number of iterations ('generations'). Defaults to 300. Using default settings may not be a good idea.

crossover The crossover method. Default is "onePoint"; also possible is "uniform".

prob The probability for switching a single bit. Defaults to 0.01; typically a small number.

pen a penalty function. Default is NULL (no penalty).

- repair a repair function. Default is NULL (no repairing).
- initP optional: the initial population. A logical matrix of size length(algo\$nB) times algo\$nP, or a function that creates such a matrix. If a function, it must take no arguments. If mode(mP) is not logical, then storage.mode(mP) will be tried (and a warning will be issued).

loopOF logical. Should the OF be evaluated through a loop? Defaults to TRUE.

- loopPen logical. Should the penalty function (if specified) be evaluated through a loop? Defaults to TRUE.
- loopRepair logical. Should the repair function (if specified) be evaluated through a loop? Defaults to TRUE.
- methodOF loop (the default), vectorised, snow or multicore. Setting vectorised is equivalent to having algo\$loopOF set to FALSE (and methodOF overrides loopOF). snow and multicore use functions clusterApply and mclapply, respectively. For snow, an object algo\$cl needs to be specified (see below). For multicore, optional arguments can be passed through algo\$mc.control (see below).
- cl a cluster object or the number of cores. See documentation of package parallel.

mc.control a list of named elements; optional settings for mclapply (for instance,

list(mc.set.seed = FALSE))

printDetail If TRUE (the default), information is printed.

printBar If TRUE (the default), a txtProgressBar is printed.

- storeF If TRUE (the default), the objective function values for every solution in every generation are stored and returned as matrix Fmat.
- storeSolutions If TRUE, the solutions (ie, binary strings) in every generation are stored and returned as a list P in list xlist (see Value section below). To check, for instance, the solutions at the end of the ith generation, retrieve xlist[[c(1L, i)]]. This will be a matrix of size algo\$nB times algo\$nP.
- classify Logical; default is FALSE. If TRUE, the result will have a class attribute TAopt attached. This feature is **experimental**: the supported methods may change without warning.

Value

A list:

xbest	the solution (the best member of the population)
OFvalue	objective function value of best solution
popF	a vector. The objective function values in the final population.
Fmat	if algo\$storeF is TRUE, a matrix of size algo\$nG times algo\$nP containing the objective function values of all solutions over the generations; else NA
xlist	if algo\$storeSolutions is TRUE, a list that contains a list P of matrices and a matrix initP (the initial solution); else NA.
initial.state	the value of .Random. seed when the function was called.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

DEopt, PSopt

greedySearch

Examples

```
## a *very* simple problem (why?):
## match a binary (logical) string y
size <- 20L ### the length of the string
OF <- function(x, y) sum(x != y)
y <- runif(size) > 0.5
x <- runif(size) > 0.5
             ### the optimum value is zero
OF(y, y)
OF(x, y)
algo <- list(nB = size, nP = 20L, nG = 100L, prob = 0.002)
sol <- GAopt(OF, algo = algo, y = y)</pre>
## show differences (if any: marked by a '^')
cat(as.integer(y), "\n", as.integer(sol$xbest), "\n",
    ifelse(y == sol$xbest , " ", "^"), "\n", sep = "")
algo$nP <- 3L ### that shouldn't work so well
sol2 <- GAopt(OF, algo = algo, y = y)</pre>
## show differences (if any: marked by a '^')
cat(as.integer(y), "\n", as.integer(sol2$xbest), "\n",
    ifelse(y == sol2$xbest , " ", "^"), "\n", sep = "")
```

greedySearch Greedy Search

Description

Greedy Search

Usage

greedySearch(OF, algo, ...)

Arguments

OF	The objective function, to be minimised. Its first argument needs to be a solu-
	tion; arguments are also passed.
algo	List of settings. See Details.
	Other variables to be passed to the objective function and to the neighbourhood function. See Details.

Details

A greedy search works starts at a provided initial solution (called the current solution) and searches a defined neighbourhood for the best possible solution. If this best neighbour is not better than the current solution, the search stops. Otherwise, the best neighbour becomes the current solution, and the search is repeated.

Value

A list:

xbest	best solution found.
OFvalue	objective function value associated with best solution.
	a matrix with two columns. <code>Fmat[,1L]</code> contains the proposed solution over all iterations; <code>Fmat[,2L]</code> contains the accepted solutions.
xlist	a list
initial.state	the value of .Random. seed when the function was called.
x0	the initial solution
iterations	the number of iterations after which the search stopped

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

See Also

LSopt

Examples

```
na <- 100
inc <- 5
R <- randomReturns(na = na,</pre>
                     ns = 1000,
                     sd = seq(0.01, 0.02, length.out = 100),
                     rho = 0.5)
S \leq cov(R)
OF <- function(x, S, ...) {
    w < -1/sum(x)
    sum(w * w * S[x, x])
}
x <- logical(na)</pre>
x[1:inc] <- TRUE</pre>
all.neighbours <- function(x, ...) {</pre>
    true <- which( x)</pre>
    false <- which(!x)</pre>
```

gridSearch

```
ans <- list()</pre>
    for (i in true) {
        for (j in false) {
             ans1 <- x
             ans1[i] <- !x[i]
             ans1[j] <- !x[j]
             ans <- c(ans, list(ans1))</pre>
        }
    }
    ans
}
algo <- list(loopOF = TRUE,</pre>
              maxit = 1000,
              all.neighbours = all.neighbours,
              x0 = x)
system.time(sol.gs <- greedySearch(OF, algo = algo, S = S))</pre>
sqrt(sol.gs$0Fvalue)
```

gridSearch

Grid Search

Description

Evaluate a function for a given list of arguments.

Usage

Arguments

fun	a function of the form $fun(x,)$, with x being a numeric vector or a list
levels	a list of levels for the arguments (see Examples)
	objects passed to fun
lower	a numeric vector. Ignored if levels are explicitly specified.
upper	a numeric vector. Ignored if levels are explicitly specified.
npar	the number of parameters. Must be supplied if lower and upper are to be ex- panded; see Details. Ignored when levels are explicitly specified, or when lower/upper are used and at least one has length greater than one. See Exam- ples.
n	the number of levels. Default is 5. Ignored if levels are explicitly specified.

printDetail	print information on the number of objective function evaluations
method	can be loop (the default), multicore or snow. See Details.
mc.control	a list containing settings that will be passed to mclapply if method is multicore. Must be a list of named elements; see the documentation of mclapply in paral- lel .
cl	default is NULL. If method snow is used, this must be a cluster object or an integer (the number of cores).
keepNames	logical: should the names of levels be kept?
asList	does fun expect a list? Default is FALSE.

Details

A grid search can be used to find 'good' parameter values for a function. In principle, a grid search has an obvious deficiency: as the length of x (the first argument to fun) increases, the number of necessary function evaluations grows exponentially. Note that gridSearch will not warn about an unreasonable number of function evaluations, but if printDetail is TRUE it will print the required number of function evaluations.

In practice, grid search is often better than its reputation. If a function takes only a few parameters, it is often a reasonable approach to find 'good' parameter values.

The function uses the mechanism of expand.grid to create the list of parameter combinations for which fun is evaluated; it calls lapply to evaluate fun if method == "loop" (the default).

If method is multicore, then function mclapply from package **parallel** is used. Further settings for mclapply can be passed through the list mc.control. If multicore is chosen but the functionality is not available, then method will be set to loop and a warning is issued. If method == "snow", the function clusterApply from package **parallel** is used. In this case, the argument cl must either be a cluster object (see the documentation of clusterApply) or an integer. If an integer, a cluster will be set up via makeCluster(c(rep("localhost", cl)), type = "SOCK") (and stopCluster is called when the function is exited). If snow is chosen but not available or cl is not specified, then method will be set to loop and a warning is issued.

Value

A list.

minfun	the minimum of fun.
minlevels	the levels that give this minimum.
values	a list. All the function values of fun.
levels	a list. All the levels for which fun was evaluated.

Author(s)

Enrico Schumann
gridSearch

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

```
testFun <- function(x)</pre>
    x[1L] + x[2L]^2
sol <- gridSearch(fun = testFun, levels = list(1:2, c(2, 3, 5)))</pre>
sol$minfun
sol$minlevels
## specify all levels
levels <- list(a = 1:2, b = 1:3)</pre>
res <- gridSearch(testFun, levels)</pre>
res$minfun
sol$minlevels
## specify lower, upper and npar
lower <- 1; upper <- 3; npar <- 2
res <- gridSearch(testFun, lower = lower, upper = upper, npar = npar)</pre>
res$minfun
sol$minlevels
## specify lower, upper, npar and n
lower <- 1; upper <- 3; npar <- 2; n <- 4
res <- gridSearch(testFun, lower = lower, upper = upper, npar = npar, n = n)</pre>
res$minfun
sol$minlevels
## specify lower, upper and n
lower <- c(1,1); upper <- c(3,3); n <- 4
res <- gridSearch(testFun, lower = lower, upper = upper, n = n)</pre>
res$minfun
sol$minlevels
## specify lower, upper (auto-expanded) and n
lower <- c(1,1); upper <- 3; n <- 4
res <- gridSearch(testFun, lower = lower, upper = upper, n = n)</pre>
res$minfun
sol$minlevels
## non-numeric inputs
test_fun <- function(x) {</pre>
    -(length(x$S) + x$N1 + x$N2)
}
```

LS.info

Local-Search Information

Description

The function can be called from the objective and neighbourhood function during a run of LSopt; it provides information such as the current iteration.

Usage

LS.info(n = 0L)

Arguments

n

generational offset; see Details.

Details

This function is still experimental.

The function can be called in the neighbourhood function or the objective function during a run of LSopt. It evaluates to a list with the state of the optimisation run, such as the current iteration.

LS. info relies on parent.frame to retrieve its information. If the function is called within another function in the neighbourhood or objective function, the argument n needs to be increased.

Value

A list iteration

iteration	current iteration
step	same as 'iteration'

LSopt

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

LSopt, TA. info

Examples

MINIMAL EXAMPLE for LSopt

ignore <- LSopt(fun, algo)</pre>

LSopt

Stochastic Local Search

Description

Performs a simple stochastic Local Search.

Usage

LSopt(OF, algo = list(), ...)

Arguments

OF	The objective function, to be minimised. Its first argument needs to be a solution; arguments are also passed.
algo	List of settings. See Details.
	Other variables to be passed to the objective function and to the neighbourhood function. See Details.

Details

Local Search (LS) changes an initial solution for a number of times, accepting only such changes that lead to an improvement in solution quality (as measured by the objective function OF). More specifically, in each iteration, a current solution xc is changed through a function algo\$neighbour. This function takes xc as an argument and returns a new solution xn. If xn is not worse than xc, ie, if OF(xn, ...) <= OF(xc, ...), then xn replaces xc.

The list algo contains the following items:

- nS The number of steps. The default is 1000; but this setting depends very much on the problem.
- nI Total number of iterations, with default NULL. If specified, it will override nS. The option is provided to makes it easier to compare and switch between functions LSopt, TAopt and SAopt.
- x0 The initial solution. This can be a function; it will then be called once without arguments to compute an initial solution, ie, x0 <- algo\$x0(). This can be useful when LSopt is called in a loop of restarts and each restart is to have its own starting value.</p>
- neighbour The neighbourhood function, called as neighbour(x, ...). Its first argument must be a solution x; it must return a changed solution.
- printDetail If TRUE (the default), information is printed. If an integer i greater then one, information is printed at very ith step.
- printBar If TRUE (the default), a txtProgressBar (from package **utils**) is printed). The progress bar is not shown if printDetail is an integer greater than 1.
- storeF if TRUE (the default), the objective function values for every solution in every generation are stored and returned as matrix Fmat.
- storeSolutions default is FALSE. If TRUE, the solutions (ie, decision variables) in every generation
 are stored and returned in list xlist (see Value section below). To check, for instance, the
 current solution at the end of the ith generation, retrieve xlist[[c(2L, i)]].
- OF.target Numeric; when specified, the algorithm will stop when an objective-function value as low as OF.target (or lower) is achieved. This is useful when an optimal objective-function value is known: the algorithm will then stop and not waste time searching for a better solution.

At the minimum, algo needs to contain an initial solution x0 and a neighbour function.

LS works on solutions through the functions neighbour and OF, which are specified by the user. Thus, a solution need not be a numeric vector, but can be any other data structure as well (eg, a list or a matrix).

To run silently (except for warnings and errors), algo\$printDetail and algo\$printBar must be FALSE.

LSopt

Value

A list:

xbest	best solution found.
OFvalue	objective function value associated with best solution.
Fmat	a matrix with two columns. Fmat[,1L] contains the proposed solution over all iterations; Fmat[,2L] contains the accepted solutions.
xlist	if algo $\$ toreSolutions is TRUE, a list; else NA. Contains the neighbour solutions at a given iteration (xn) and the current solutions (xc). Example: Fmat[i, 2L] is the objective function value associated with xlist[[c(2L, i)]].
initial.state	the value of .Random.seed when the function was called.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). <code>https://enricoschumann.net/NMOF.htm#NMOFmanual</code>

See Also

TAopt, restartOpt. Package **neighbours** (also on CRAN) offers helpers for creating neighbourhood functions.

Examples

Aim: find the columns of X that, when summed, give y

```
## random data set
nc <- 25L
                    ## number of columns in data set
                    ## number of rows in data set
nr <- 5L
howManyCols <- 5L ## length of true solution</pre>
X <- array(runif(nr*nc), dim = c(nr, nc))</pre>
xTRUE <- sample(1L:nc, howManyCols)</pre>
Xt <- X[ , xTRUE, drop = FALSE]</pre>
y <- rowSums(Xt)</pre>
## a random solution x0 ...
makeRandomSol <- function(nc) {</pre>
    ii <- sample.int(nc, sample.int(nc, 1L))</pre>
    x0 <- logical(nc); x0[ii] <- TRUE</pre>
    x0
}
x0 <- makeRandomSol(nc)</pre>
```

```
## ... but probably not a good one
sum(y - rowSums(X[ , xTRUE, drop = FALSE])) ## should be 0
sum(y - rowSums(X[ , x0, drop = FALSE]))
## a neighbourhood function: switch n elements in solution
neighbour <- function(xc, Data) {</pre>
    xn <- xc
    p <- sample.int(Data$nc, Data$n)</pre>
    xn[p] <- !xn[p]</pre>
    if (sum(xn) < 1L)
        xn <- xc
    xn
}
## a greedy neighbourhood function
neighbourG <- function(xc, Data) {</pre>
    of <- function(x)
        abs(sum(Data$y - rowSums(Data$X[ ,x, drop = FALSE])))
    xbest <- xc
    Fxbest <- of(xbest)</pre>
    for (i in 1L:Data$nc) {
        xn <- xc; p <- i
        xn[p] <- !xn[p]</pre>
        if (sum(xn) >= 1L) {
            Fxn <- of(xn)</pre>
            if (Fxn < Fxbest) {</pre>
                xbest <- xn
                 Fxbest <- Fxn
            }
        }
    }
    xbest
}
## an objective function
OF <- function(xn, Data)
    abs(sum(Data$y - rowSums(Data$X[ ,xn, drop = FALSE])))
## (1) GREEDY SEARCH
## note: this could be done in a simpler fashion, but the
##
         redundancies/overhead here are small, and the example is to
         show how LSopt can be used for such a search
##
Data <- list(X = X, y = y, nc = nc, nr = nr, n = 1L)
algo <- list(nS = 500L, neighbour = neighbourG, x0 = x0,
             printBar = FALSE, printDetail = FALSE)
solG <- LSopt(OF, algo = algo, Data = Data)</pre>
## after how many iterations did we stop?
iterG <- min(which(solG$Fmat[ ,2L] == solG$OFvalue))</pre>
solG$OFvalue ## the true solution has OF-value 0
## (2) LOCAL SEARCH
```

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MA

```
algo$neighbour <- neighbour
solLS <- LSopt(OF, algo = algo, Data = Data)</pre>
iterLS <- min(which(solLS$Fmat[ ,2L] == solLS$OFvalue))</pre>
solLS$OFvalue ## the true solution has OF-value 0
## (3) *Threshold Accepting*
algo$nT <- 10L
algo$nS <- ceiling(algo$nS/algo$nT)</pre>
algo$q <- 0.99
solTA <- TAopt(OF, algo = algo, Data = Data)</pre>
iterTA <- min(which(solTA$Fmat[ ,2L] == solTA$OFvalue))</pre>
solTA$OFvalue ## the true solution has OF-value 0
## look at the solution
all <- sort(unique(c(which(solTA$xbest),</pre>
                     which(solLS$xbest),
                     which(solG$xbest),
                     xTRUE)))
ta <- ls <- greedy <- true <- character(length(all))</pre>
true[ match(xTRUE, all)] <- "o"</pre>
greedy[match(which(solG$xbest), all)] <- "o"</pre>
       match(which(solLS$xbest), all)] <- "o"</pre>
lsΓ
       match(which(solTA$xbest), all)] <- "o"</pre>
ta[
data.frame(true = true, greedy = greedy, LS = ls , TA = ta,
           row.names=all)
## plot results
par(ylog = TRUE, mar = c(5,5,1,6), las = 1)
plot(solTA$Fmat[seq_len(iterTA) ,2L],type = "1", log = "y",
     ylim = c(1e-4),
              max(pretty(c(solG$Fmat,solLS$Fmat,solTA$Fmat)))),
     xlab = "iterations", ylab = "OF value", col = grey(0.5))
lines(cummin(solTA$Fmat[seq_len(iterTA), 2L]), type = "1")
lines(solG$Fmat[ seq_len(iterG), 2L], type = "p", col = "blue")
lines(solLS$Fmat[seq_len(iterLS), 2L], type = "1", col = "goldenrod3")
legend(x = "bottomleft",
       legend = c("TA best solution", "TA current solution",
                   "Greedy", "LS current/best solution"),
       lty = c(1,1,0,1),
       col = c("black",grey(0.5),"blue","goldenrod2"),
       pch = c(NA, NA, 21, NA))
axis(4, at = c(solG$0Fvalue, solLS$0Fvalue, solTA$0Fvalue),
        labels = NULL, las = 1)
lines(x = c(iterG, par()$usr[2L]), y = rep(solG$0Fvalue,2),
      col = "blue", lty = 3)
lines(x = c(iterTA, par()$usr[2L]), y = rep(solTA$0Fvalue,2),
      col = "black", lty = 3)
lines(x = c(iterLS, par()$usr[2L]), y = rep(solLS$0Fvalue,2),
      col = "goldenrod3", lty = 3)
```

Description

The function computes a moving average of a vector.

Usage

MA(y, order, pad = NULL)

Arguments

У	a numeric vector
order	An integer. The order of the moving average. The function is defined such that order one returns y (see Examples).
pad	Defaults to NULL. If not NULL, all elements of the returned moving average with position smaller than order are replaced by the value of pad. Sensible values may be NA or 0.

Value

Returns a vector of length length(y).

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

```
MA(1:10, 3)
MA(1:10, 3, pad = NA)
y \le seq(1, 4, by = 0.3)
z <- MA(y, 1)
all(y == z)
                 ### (typically) FALSE
all.equal(y, z) ### should be TRUE
## 'Relative strength index'
rsi <- function(y, t) {</pre>
    y \leq -diff(y)
    ups <- y + abs(y)
    downs <- y - abs(y)
    RS <- -MA(ups, t) / MA(downs, t)
    RS/(1 + RS)
}
x <- cumprod(c(100, 1 + rnorm(100, sd = 0.01)))</pre>
```

maxSharpe

```
par(mfrow = c(2,1))
plot(x, type = "l")
plot(rsi(x, 14), ylim = c(0,1), type = "l")
```

maxSharpe

Maximum-Sharpe-Ratio/Tangency Portfolio

Description

Compute maximum Sharpe-ratio portfolios, subject to lower and upper bounds on weights.

Usage

```
maxSharpe(m, var, min.return,
    wmin = -Inf, wmax = Inf, method = "qp",
    groups = NULL, groups.wmin = NULL, groups.wmax = NULL)
```

Arguments

m	vector of expected (excess) returns.
var	the covariance matrix: a numeric (real), symmetric matrix
min.return	minimumm required return. This is a technical parameter, used only for QP.
wmin	numeric: a lower bound on weights. May also be a vector that holds specific bounds for each asset.
wmax	numeric: an upper bound on weights. May also be a vector that holds specific bounds for each asset.
method	character. Currently, only "qp" is supported.
groups	a list of group definitions
groups.wmin	a numeric vector
groups.wmax	a numeric vector

Details

The function uses solve.QP from package **quadprog**. Because of the algorithm that solve.QP uses, var has to be positive definit (i.e. must be of full rank).

Value

a numeric vector (the portfolio weights) with an attribute variance (the portfolio's variance)

Author(s)

Enrico Schumann

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Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Schumann, E. (2012) Computing the global minimum-variance portfolio. https://enricoschumann. net/R/minvar.htm

See Also

minvar, mvPortfolio, mvFrontier

Examples

```
S <- var(R <- NMOF::randomReturns(3, 10, 0.03))
x <- solve(S, colMeans(R))
x/sum(x)
x <- coef(lm(rep(1, 10) ~ -1 + R))
unname(x/sum(x))
maxSharpe(m = colMeans(R), var = S)
maxSharpe(m = colMeans(R), var = S, wmin = 0, wmax = 1)</pre>
```

|--|

Option Pricing via Monte-Carlo Simulation

Description

Functions to calculate the theoretical prices of options through simulation.

Usage

```
gbm(npaths, timesteps, r, v, tau, S0,
    exp.result = TRUE, antithetic = FALSE)
gbb(npaths, timesteps, S0, ST, v, tau,
    log = FALSE, exp.result = TRUE)
```

Arguments

npaths	the number of paths
timesteps	timesteps per path
r	the mean per unit of time
v	the variance per unit of time
tau	time
S0	initial value

ST	final value of Brownian bridge
log	logical: construct bridge from log series?
exp.result	logical: compute exp of the final path, or return log values?
antithetic	logical: if TRUE, random numbers for only npaths/2 are drawn, and the random numbers are mirrored

Details

gbm generates sample paths of geometric Brownian motion.

gbb generates sample paths of a Brownian bridge by first creating paths of Brownian motion W from time 0 to time T, with W_0 equal to zero. Then, at each t, it subtracts $t/T * W_T$ and adds S0*(1-t/T)+ST*(t/T).

Value

A matrix of sample paths; each column contains the price path of an asset. Even with only a single time-step, the matrix will have two rows (the first row is S0).

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

vanillaOptionEuropean

```
## price a European option
## ... parameters
npaths <- 5000 ## increase number to get more precise results
timesteps <- 1
S0 <- 100
ST <- 100
tau <- 1
r <- 0.01
v <- 0.25^2
## ... create paths
paths <- gbm(npaths, timesteps, r, v, tau, S0 = S0)
## ... a helper function</pre>
```

```
mc <- function(paths, payoff, ...)</pre>
    payoff(paths, ...)
## ... a payoff function (European call)
payoff <- function(paths, X, r, tau)</pre>
    exp(-r * tau) * mean(pmax(paths[NROW(paths), ] - X, 0))
## ... compute and check
mc(paths, payoff, X = 100, r = r, tau = tau)
vanillaOptionEuropean(S0, X = 100, tau = tau, r = r, v = v)$value
## compute delta via forward difference
## (see Gilli/Maringer/Schumann, ch. 9)
h <- 1e-6
                           ## a small number
rnorm(1)
                           ## make sure RNG is initialised
rnd.seed <- .Random.seed ## store current seed</pre>
paths1 <- gbm(npaths, timesteps, r, v, tau, S0 = S0)</pre>
.Random.seed <- rnd.seed
paths2 <- gbm(npaths, timesteps, r, v, tau, S0 = S0 + h)</pre>
delta.mc <- (mc(paths2, payoff, X = 100, r = r, tau = tau)-</pre>
             mc(paths1, payoff, X = 100, r = r, tau = tau))/h
delta <- vanillaOptionEuropean(S0, X = 100, tau = tau,</pre>
                                r = r, v = v)$delta
delta.mc - delta
## a fanplot
steps <- 100
paths <- results <- gbm(1000, steps, r = 0, v = 0.2^2,</pre>
                         tau = 1, S0 = 100)
levels <- seq(0.01, 0.49, length.out = 20)</pre>
greys <- seq(0.9, 0.50, length.out = length(levels))</pre>
## start with an empty plot ...
plot(0:steps, rep(100, steps+1), ylim = range(paths),
     xlab = "", ylab = "", lty = 0, type = "1")
## ... and add polygons
for (level in levels) {
    l <- apply(paths, 1, quantile, level)</pre>
    u <- apply(paths, 1, quantile, 1 - level)</pre>
    col <- grey(greys[level == levels])</pre>
    polygon(c(0:steps, steps:0), c(l, rev(u)),
            col = col, border = NA)
    ## add border lines
    ## lines(0:steps, 1, col = grey(0.4))
```

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minCVaR

```
## lines(0:steps, u, col = grey(0.4))
}
```

minCVaR

Minimum Conditional-Value-at-Risk (CVaR) Portfolios

Description

Compute minimum-CVaR portfolios, subject to lower and upper bounds on weights.

Usage

```
minCVaR(R, q = 0.1, wmin = 0, wmax = 1,
    min.return = NULL, m = NULL,
    method = "Rglpk",
    groups = NULL, groups.wmin = NULL, groups.wmax = NULL,
    Rglpk.control = list())
```

Arguments

R	the scenario matrix: a numeric (real) matrix
q	the Value-at-Risk level: a number between 0 and 0.5
wmin	numeric: a lower bound on weights. May also be a vector that holds specific bounds for each asset.
wmax	numeric: an upper bound on weights. May also be a vector that holds specific bounds for each asset.
m	vector of expected returns. Only used if min.return is specified.
min.return	minimal required return. If m is not specified, the column means of R are used.
method	character. Currently, only "Rglpk" is supported.
groups	a list of group definitions
groups.wmin	a numeric vector
groups.wmax	a numeric vector
Rglpk.control	a list: settings passed to Rglpk_solve_LP

Details

Compute the minimum CVaR portfolio for a given scenario set. The default method uses the formulation as a Linear Programme, as described in Rockafellar/Uryasev (2000).

The function uses Rglpk_solve_LP from package **Rglpk**.

Value

a numeric vector (the portfolio weights); attached is an attribute whose name matches the method name

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Rockafellar, R. T. and Uryasev, S. (2000). Optimization of Conditional Value-at-Risk. Journal of Risk. 2 (3), 21–41.

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

Schumann, E. (2020) Minimising Conditional Value-at-Risk (CVaR). https://enricoschumann. net/notes/minimising-conditional-var.html

See Also

minvar

Examples

```
if (requireNamespace("Rglpk")) {
   ns <- 5000 ## number of scenarios
   na <- 20 ## number of assets
   R <- randomReturns(na, ns, sd = 0.01, rho = 0.5)
   res <- minCVaR(R, 0.25)
   c(res) ## portfolio weights
}</pre>
```

minMAD

Compute Minimum Mean–Absolute-Deviation Portfolios

Description

Compute minimum mean-absolute-deviation portfolios.

Usage

```
minMAD(R, wmin = 0, wmax = 1,
    min.return = NULL, m = NULL, demean = TRUE,
    method = "lp",
    groups = NULL, groups.wmin = NULL, groups.wmax = NULL,
    Rglpk.control = list())
```

minMAD

Arguments

R	a matrix of return scenarios: each column represents one asset; each row repre- sents one scenario
wmin	minimum weight
wmax	maximum weight
min.return	a minimum required return; ignored if NULL
m	a vector of expected returns. If NULL, but min.return is not NULL, then column means are used as expected returns.
demean	logical. If TRUE, the columns of R are demeaned, corresponding to an objective function xxxx
method	string. Supported are lp and ls.
groups	group definitions
groups.wmin	list of vectors
groups.wmax	list of vectors
Rglpk.control	a list

Details

Compute the minimum mean-absolute-deviation portfolio for a given scenario set.

The function uses Rglpk_solve_LP from package **Rglpk**.

Value

a vector of portfolio weights

Author(s)

Enrico Schumann

References

Konno, H. and Yamazaki, H. (1991) Mean-Absolute Deviation Portfolio Optimization Model and Its Applications to Tokyo Stock Market. *Management Science*. **37** (5), 519–531.

See Also

minvar,minCVaR

minvar

Description

Compute minimum-variance portfolios, subject to lower and upper bounds on weights.

Usage

```
minvar(var, wmin = 0, wmax = 1, method = "qp",
    groups = NULL, groups.wmin = NULL, groups.wmax = NULL)
```

Arguments

var	the covariance matrix: a numeric (real), symmetric matrix
wmin	numeric: a lower bound on weights. May also be a vector that holds specific bounds for each asset.
wmax	numeric: an upper bound on weights. May also be a vector that holds specific bounds for each asset.
method	character. Currently, only "qp" is supported.
groups	a list of group definitions
groups.wmin	a numeric vector
groups.wmax	a numeric vector

Details

For method "qp", the function uses solve.QP from package **quadprog**. Because of the algorithm that solve.QP uses, var has to be positive definite (i.e. must be of full rank).

Value

a numeric vector (the portfolio weights) with an attribute variance (the portfolio's variance)

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

Schumann, E. (2012) Computing the global minimum-variance portfolio. https://enricoschumann. net/R/minvar.htm

minvar

See Also

TAopt

```
## variance-covariance matrix from daily returns, 1 Jan 2014 -- 31 Dec 2013, of
## cleaned data set at https://enricoschumann.net/data/gilli_accuracy.html
```

```
if (requireNamespace("quadprog")) {
```

```
var <- structure(c(0.000988087100677907, -0.0000179669410403153, 0.000368923882626859,
                 0.000208303611101873, 0.000262742052359594, -0.0000179669410403153,
                  0.00171852167358765, 0.0000857467457561209, 0.0000215059246610556,
                  0.0000283532159921211, 0.000368923882626859, 0.0000857467457561209,
                   0.00075871953281751, 0.000194002299424151, 0.000188824454515841,
                  0.000208303611101873, 0.0000215059246610556, 0.000194002299424151,
                   0.000265780633005374,\ 0.000132611196599808,\ 0.000262742052359594,
                  0.0000283532159921211, 0.000188824454515841, 0.000132611196599808,
                   0.00025948420130626),
                  .Dim = c(5L, 5L),
               .Dimnames = list(c("CBK.DE", "VOW.DE", "CON.DE", "LIN.DE", "MUV2.DE"),
                               c("CBK.DE", "VOW.DE", "CON.DE", "LIN.DE", "MUV2.DE")))
 ##
              CBK.DE
                         VOW.DE
                                   CON.DE
                                           LIN.DE MUV2.DE
 ## CBK.DE 0.000988 -0.0000180 0.0003689 0.0002083 0.0002627
 ## VOW.DE -0.000018 0.0017185 0.0000857 0.0000215 0.0000284
 ## CON.DE 0.000369 0.0000857 0.0007587 0.0001940 0.0001888
 ## LIN.DE 0.000208 0.0000215 0.0001940 0.0002658 0.0001326
 ## MUV2.DE 0.000263 0.0000284 0.0001888 0.0001326 0.0002595
 ##
minvar(var, wmin = 0, wmax = 0.5)
minvar(var,
       wmin = c(0.1,0,0,0,0), ## enforce at least 10% weight in CBK.DE
       wmax = 0.5)
minvar(var, wmin = -Inf, wmax = Inf)
                                       ## no bounds
 ## [1] -0.0467 0.0900 0.0117 0.4534 0.4916
minvar(var, wmin = -Inf, wmax = 0.45) ## no lower bounds
 ## [1] -0.0284 0.0977 0.0307 0.4500 0.4500
minvar(var, wmin = 0.1, wmax = Inf)
                                       ## no upper bounds
 ## [1] 0.100 0.100 0.100 0.363 0.337
 ## group constraints:
     group 1 consists of asset 1 only, and must have weight [0.25,0.30]
 ##
    group 2 consists of assets 4 and 5, and must have weight [0.10,0.20]
 ##
 ## => unconstrained
minvar(var, wmin = 0, wmax = 0.40)
```

```
## [1] 0.0097 0.1149 0.0754 0.4000 0.4000
## => with group constraints
minvar(var, wmin = 0, wmax = 0.40,
    groups = list(1, 4:5),
    groups.wmin = c(0.25, 0.1),
    groups.wmax = c(0.30, 0.2))
## [1] 0.250 0.217 0.333 0.149 0.051
}
```

mvFrontier

Computing Mean–Variance Efficient Portfolios

Description

Compute mean-variance efficient portfolios and efficient frontiers.

Usage

```
mvFrontier(m, var, wmin = 0, wmax = 1, n = 50, rf = NA,
            groups = NULL, groups.wmin = NULL, groups.wmax = NULL)
mvPortfolio(m, var, min.return, wmin = 0, wmax = 1, lambda = NULL,
            groups = NULL, groups.wmin = NULL, groups.wmax = NULL)
```

Arguments

m	vector of expected returns
var	expected variance-covariance matrix
wmin	numeric: minimum weights
wmax	numeric: maximum weights
n	number of points on the efficient frontier
min.return	minimal required return
rf	risk-free rate
lambda	risk-reward trade-off
groups	a list of group definitions
groups.wmin	a numeric vector
groups.wmax	a numeric vector

Details

mvPortfolio computes a single mean-variance efficient portfolio, using package **quadprog**. It does so by minimising portfolio variance, subject to constraints on minimum return and budget (weights need to sum to one), and min/max constraints on the weights.

mvFrontier

If λ is specified, the function ignores the min. return constraint and instead solves the model

$$\min_{w} -\lambda \mathsf{m}' w + (1-\lambda) w' \mathsf{var} w$$

in which w are the weights. If λ is a vector of length 2, then the model becomes

$$\min_{w} \ -\lambda_1 \mathsf{m}' w + \lambda_2 w' \mathsf{var} w$$

which may be more convenient (e.g. for setting λ_1 to 1).

mvFrontier computes returns, volatilities and compositions for portfolios along an efficient frontier. If rf is not NA, cash is included as an asset.

Value

For mvPortfolio, a numeric vector of weights.

For mvFrontier, a list of three components:

return	returns of portfolios
volatility	volatilities of portfolios
weights	A matrix of portfolio weights. Each column holds the weights for one portfolio on the frontier. If rf is specified, an additional row is added, providing the cash weight.

The *i*-th portfolio on the frontier corresponds to the *i*-th elements of return and volatility, and the *i*-th column of portfolio.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

minvar for computing the minimum-variance portfolio

```
na <- 4
vols <- c(0.10, 0.15, 0.20,0.22)
m <- c(0.06, 0.12, 0.09, 0.07)
const_cor <- function(rho, na) {
    C <- array(rho, dim = c(na, na))
    diag(C) <- 1</pre>
```

```
С
}
var <- diag(vols) %*% const_cor(0.5, na) %*% diag(vols)</pre>
wmax <- 1
                   # maximum holding size
wmin <- 0.0
                     # minimum holding size
rf <- 0.02
if (requireNamespace("quadprog")) {
  p1 <- mvFrontier(m, var, wmin = wmin, wmax = wmax, n = 50)</pre>
  p2 <- mvFrontier(m, var, wmin = wmin, wmax = wmax, n = 50, rf = rf)</pre>
  plot(p1$volatility, p1$return, pch = 19, cex = 0.5, type = "o",
       xlab = "Expected volatility",
       ylab = "Expected return")
  lines(p2$volatility, p2$return, col = grey(0.5))
  abline(v = 0, h = rf)
} else
  message("Package 'quadprog' is required")
```

```
NS
```

Zero Rates for Nelson-Siegel-Svensson Model

Description

Compute zero yields for Nelson-Siegel (NS)/Nelson-Siegel-Svensson (NSS) model.

Usage

NS(param, tm) NSS(param, tm)

Arguments

param	a vector. For NS: β_1 , β_2 , β_3 , λ . For NSS: a vector: β_1 , β_2 , β_3 , β_4 , λ_1 , λ_2
tm	a vector of maturities

Details

See Chapter 14 in Gilli/Maringer/Schumann (2011). Maturities (tm) need to be given in time (not dates).

Value

The function returns a vector of length length(tm).

Author(s)

Enrico Schumann

References

Gilli, M. and Grosse, S. and Schumann, E. (2010) Calibrating the Nelson-Siegel-Svensson model, COMISEF Working Paper Series No. 031. https://enricoschumann.net/COMISEF/wps031.pdf

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Gilli, M. and Schumann, E. (2010) A Note on 'Good' Starting Values in Numerical Optimisation, COMISEF Working Paper Series No. 044. https://enricoschumann.net/COMISEF/wps044.pdf

Nelson, C.R. and Siegel, A.F. (1987) Parsimonious Modeling of Yield Curves. *Journal of Business*, **60**(4), pp. 473–489.

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Svensson, L.E. (1994) Estimating and Interpreting Forward Interest Rates: Sweden 1992–1994. IMF Working Paper 94/114.

See Also

NSf, NSSf

Examples

NSf

Factor Loadings for Nelson-Siegel and Nelson-Siegel-Svensson

Description

Computes the factor loadings for Nelson–Siegel (NS) and Nelson–Siegel–Svensson (NSS) model for given lambda values.

Usage

NSf(lambda, tm) NSSf(lambda1, lambda2, tm)

NSf

Arguments

lambda	the λ parameter of the NS model (a scalar)
lambda1	the λ_1 parameter of the NSS model (a scalar)
lambda2	the λ_2 parameter of the NSS model (a scalar)
tm	a numeric vector with times-to-payment/maturity

Details

The function computes the factor loadings for given λ parameters. Checking the correlation between these factor loadings can help to set reasonable λ values for the NS/NSS models.

Value

For NS, a matrix with length(tm) rows and three columns. For NSS, a matrix with length(tm) rows and four columns.

Author(s)

Enrico Schumann

References

Gilli, M. and Grosse, S. and Schumann, E. (2010) Calibrating the Nelson-Siegel-Svensson model, COMISEF Working Paper Series No. 031. https://enricoschumann.net/COMISEF/wps031.pdf

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Gilli, M. and Schumann, E. (2010) A Note on 'Good' Starting Values in Numerical Optimisation, COMISEF Working Paper Series No. 044. https://enricoschumann.net/COMISEF/wps044.pdf

Nelson, C.R. and Siegel, A.F. (1987) Parsimonious Modeling of Yield Curves. *Journal of Business*, **60**(4), pp. 473–489.

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Svensson, L.E. (1994) Estimating and Interpreting Forward Interest Rates: Sweden 1992–1994. IMF Working Paper 94/114.

See Also

NS, NSS

```
## Nelson-Siegel
cor(NSf(lambda = 6, tm = 1:10)[-1L, -1L])
## Nelson-Siegel-Svensson
cor(NSSf(lambda1 = 1, lambda2 = 5, tm = 1:10)[-1L, -1L])
cor(NSSf(lambda1 = 4, lambda2 = 9, tm = 1:10)[-1L, -1L])
```

optionData

Description

Closing prices of DAX index options as of 2012-02-10.

Usage

optionData

Format

optionData is a list with six components:

pricesCall a matrix of size 124 times 10. The rows are the strikes; each column belongs to one expiry date.

pricesPut a matrix of size 124 times 10

index The DAX index (spot).

future The available future settlement prices.

Euribor Euribor rates.

NSSpar Paramaters for German government bond yields, as estimated by the Bundesbank.

Details

Settlement prices for EUREX options are computed at 17:30, Frankfurt Time, even though trading continues until 22:00.

Source

The data was obtained from several websites: close prices of EUREX products were collected from https://www.eurex.com/ex-en/; Euribor rates and the parameters of the Nelson-Siegel-Svensson can be found at https://www.bundesbank.de/en/.

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Examples

str(optionData)
NSS(optionData\$NSSpar, 1:10)

pm

Description

Compute partial moments.

Usage

```
pm(x, xp = 2, threshold = 0, lower = TRUE,
normalise = FALSE, na.rm = FALSE)
```

Arguments

х	a numeric vector or a matrix
хр	exponent
threshold	a numeric vector of length one
lower	logical
normalise	logical
na.rm	logical

Details

For a vector x of length n, partial moments are computed as follows:

upper partial moment
$$=$$
 $\frac{1}{n} \sum_{x>t} (x-t)^{\epsilon}$
lower partial moment $=$ $\frac{1}{n} \sum_{x$

The threshold is denoted t, the exponent xp is labelled e.

If normalise is TRUE, the result is raised to 1/xp. If x is a matrix, the function will compute the partial moments column-wise.

See Gilli, Maringer and Schumann (2019), chapter 14.

Value

numeric

Author(s)

Enrico Schumann

PSopt

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Examples

```
pm(x <- rnorm(100), 2)
var(x)/2
pm(x, 2, normalise = TRUE)
sqrt(var(x)/2)</pre>
```

PSopt

Particle Swarm Optimisation

Description

The function implements Particle Swarm Optimisation.

Usage

PSopt(OF, algo = list(), ...)

Arguments

OF	the objective function to be minimised. See Details.
algo	a list with the settings for algorithm. See Details and Examples.
	pieces of data required to evaluate the objective function. See Details.

Details

The function implements Particle Swarm Optimisation (PS); see the references for details on the implementation. PS is a population-based optimisation heuristic. It develops several solutions (a 'population') over a number of iterations. PS is directly applicable to continuous problems since the population is stored in real-valued vectors. In each iteration, a solution is updated by adding another vector called velocity. Think of a solution as a position in the search space, and of velocity as the direction into which this solution moves. Velocity changes over the course of the optimization: it is biased towards the best solution found by the particular solution and the best overall solution. The algorithm stops after a fixed number of iterations.

To allow for constraints, the evaluation works as follows: after a new solution is created, it is (i) repaired, (ii) evaluated through the objective function, (iii) penalised. Step (ii) is done by a call to OF; steps (i) and (iii) by calls to algo\$repair and algo\$pen. Step (i) and (iii) are optional, so the respective functions default to NULL. A penalty can also be directly written in the OF, since it amounts to a positive number added to the 'clean' objective function value. It can be advantageous

to write a separate penalty function if either only the objective function or only the penalty function can be vectorised. (Constraints can also be added without these mechanisms. Solutions that violate constraints can, for instance, be mapped to feasible solutions, but without actually changing them. See Maringer and Oyewumi, 2007, for an example with Differential Evolution.)

Conceptually, PS consists of two loops: one loop across the iterations and, in any given generation, one loop across the solutions. This is the default, controlled by the variables algo\$loopOF, algo\$loopRepair, algo\$loopPen and loopChangeV which all default to TRUE. But it does not matter in what order the solutions are evaluated (or repaired or penalised), so the second loop can be vectorised. Examples are given in the vignettes and in the book. The respective algo\$loopFun must then be set to FALSE.

The objective function, the repair function and and the penalty function will be called as fun(solution, ...).

- The list algo contains the following items:
- nP population size. Defaults to 100. Using default settings may not be a good idea.
- nG number of iterations. Defaults to 500. Using default settings may not be a good idea.
- c1 the weight towards the individual's best solution. Typically between 0 and 2; defaults to 1. Using default settings may not be a good idea. In some cases, even negative values work well: the solution is then driven off its past best position. For 'simple' problems, setting c1 to zero may work well: the population moves then towards the best overall solution.
- c2 the weight towards the populations's best solution. Typically between 0 and 2; defaults to 1. Using default settings may not be a good idea. In some cases, even negative values work well: the solution is then driven off the population's past best position.
- iner the inertia weight (a scalar), which reduces velocity. Typically between 0 and 1. Default is 0.9.
- initV the standard deviation of the initial velocities. Defaults to 1.
- maxV the maximum (absolute) velocity. Setting limits to velocity is sometimes called velocity clamping. Velocity is the change in a given solution in a given iteration. A maximum velocity can be set so to prevent unreasonable velocities ('overshooting'): for instance, if a decision variable may lie between 0 and 1, then an absolute velocity much greater than 1 makes rarely sense.
- min, max vectors of minimum and maximum parameter values. The vectors min and max are used to determine the dimension of the problem and to randomly initialise the population. Per default, they are no constraints: a solution may well be outside these limits. Only if algo\$minmaxConstr is TRUE will the algorithm repair solutions outside the min and max range.
- minmaxConstr if TRUE, algo\$min and algo\$max are considered constraints. Default is FALSE.
- pen a penalty function. Default is NULL (no penalty).
- repair a repair function. Default is NULL (no repairing).
- changeV a function to change velocity. Default is NULL (no change). This function is called before the velocity is added to the current solutions; it can be used to impose restrictions like changing only a number of decision variables.
- initP optional: the initial population. A matrix of size length(algo\$min) times algo\$nP, or a function that creates such a matrix. If a function, it should take no arguments.
- loopOF logical. Should the OF be evaluated through a loop? Defaults to TRUE.

- loopPen logical. Should the penalty function (if specified) be evaluated through a loop? Defaults to TRUE.
- loopRepair logical. Should the repair function (if specified) be evaluated through a loop? Defaults to TRUE.
- loopChangeV logical. Should the changeV function (if specified) be evaluated through a loop? Defaults to TRUE.
- printDetail If TRUE (the default), information is printed. If an integer i greater then one, information is printed at very ith iteration.
- printBar If TRUE (the default), a txtProgressBar (from package utils) is printed).
- storeF If TRUE (the default), the objective function values for every solution in every generation are stored and returned as matrix Fmat.
- storeSolutions default is FALSE. If TRUE, the solutions (ie, decision variables) in every generation are stored as lists P and Pbest, both stored in the list xlist which the function returns. To check, for instance, the solutions at the end of the ith iteration, retrieve xlist[[c(1L, i)]]; the best solutions at the end of this iteration are in xlist[[c(2L, i)]]. P[[i]] and Pbest[[i]] will be matrices of size length(algo\$min) times algo\$nP.
- classify Logical; default is FALSE. If TRUE, the result will have a class attribute TAopt attached. This feature is **experimental**: the supported methods may change without warning.
- drop Default is TRUE. If FALSE, the dimension is not dropped from a single solution when it is passed to a function. (That is, the function will receive a single-column matrix.)

Value

Returns a list:

xbest	the solution
OFvalue	objective function value of best solution
рорҒ	a vector: the objective function values in the final population
Fmat	if algo\$storeF is TRUE, a matrix of size algo\$nG times algo\$nP. Each column contains the best objective function value found by the particular solution.
xlist	if algo\$storeSolutions is TRUE, a list that contains two lists P and Pbest of matrices, and a matrix initP (the initial solution); else NA.
initial.state	the value of .Random.seed when PSopt was called.

Author(s)

Enrico Schumann

References

Eberhart, R.C. and Kennedy, J. (1995) A New Optimizer using Particle Swarm theory. *Proceedings* of the Sixth International Symposium on Micromachine and Human Science, pp. 39–43.

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

DEopt

Examples

```
## Least Median of Squares (LMS) estimation
genData <- function(nP, n0, ol, dy) {</pre>
    ## create dataset as in Salibian-Barrera & Yohai 2006
    ## nP = regressors, n0 = number of obs
    ## ol = number of outliers, dy = outlier size
    mRN <- function(m, n) array(rnorm(m * n), dim = c(m, n))</pre>
    y <- mRN(n0, 1)
    X <- cbind(as.matrix(numeric(n0) + 1), mRN(n0, nP - 1L))</pre>
    zz <- sample(n0)
    z <- cbind(1, 100, array(0, dim = c(1L, nP - 2L)))</pre>
    for (i in seq_len(ol)) {
        X[zz[i], ] <- z
        y[zz[i]] <- dy
    }
    list(X = X, y = y)
}
OF <- function(param, data) {
    X <- data$X
    y <- data$y
    aux <- as.vector(y) - X %*% param</pre>
    ## as.vector(y) for recycling (param is a matrix)
    aux <- aux * aux
    aux <- apply(aux, 2, sort, partial = data$h)</pre>
    aux[h, ]
}
nP <- 2L; nO <- 100L; ol <- 10L; dy <- 150
aux <- genData(nP,n0,ol,dy); X <- aux$X; y <- aux$y</pre>
h <- (n0 + nP + 1L) %/% 2
data <- list(y = y, X = X, h = h)
algo <- list(min = rep(-10, nP), max = rep( 10, nP),
    c1 = 1.0, c2 = 2.0,
    iner = 0.7, initV = 1, maxV = 3,
    nP = 100L, nG = 300L, loopOF = FALSE)
system.time(sol <- PSopt(OF = OF, algo = algo, data = data))</pre>
if (require("MASS", quietly = TRUE)) {
    ## for nsamp = "best", in this case, complete enumeration
    ## will be tried. See ?lqs
    system.time(test1 <- lqs(data$y ~ data$X[, -1L],</pre>
            adjust = TRUE,
            nsamp = "best",
            method = "lqs",
            quantile = data$h))
```

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putCallParity

```
}
## check
x1 <- sort((y - X %*% as.matrix(sol$xbest))^2)[h]
cat("Particle Swarm\n",x1,"\n\n")
if (require("MASS", quietly = TRUE)) {
    x2 <- sort((y - X %*% as.matrix(coef(test1)))^2)[h]
    cat("lqs\n", x2, "\n\n")
}</pre>
```

putCallParity Put-Call Parity

Description

Put-call parity

Usage

```
putCallParity(what, call, put, S, X, tau, r, q = 0, tauD = 0, D = 0)
```

Arguments

what	character: what to compute. Currently only call or put are supported.
call	call price
put	put price
S	underlier
Х	strike
tau	time to expiry
r	interest rate
q	dividend rate
tauD	numeric vector: time to dividend
D	numeric vector: dividends

Details

Put-call parity only works for European options. The function is vectorised (like vanillaOptionEuropean), except for dividends.

Value

Numeric vector.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Examples

```
S <- 100; X <- 100; tau <- 1; r <- 0.02; q <- 0.0;
vol <- 0.3; D <- 20; tauD <- 0.5
call <- vanillaOptionEuropean(S, X, tau, r, q, vol^2,</pre>
                               tauD = tauD, D = D, type = "call")$value
put <- vanillaOptionEuropean(S, X, tau, r, q, vol^2,</pre>
                               tauD = tauD, D = D, type = "put")$value
## recover the call from the put (et vice versa)
all.equal(call, putCallParity("call", put = put, S=S, X=X, tau=tau,
                              r=r, q=q, tauD=tauD, D=D))
all.equal(put, putCallParity("put", call = call, S=S, X=X, tau=tau,
                              r=r, q=q, tauD=tauD, D=D))
## Black--Scholes--Merton with with 'callCF'
S <- 100; X <- 90; tau <- 1; r <- 0.02; q <- 0.08
v <- 0.2<sup>2</sup> ## variance, not volatility
(ccf <- callCF(cf = cfBSM, S = S, X = X, tau = tau, r = r, q = q,
                 v = v, implVol = TRUE))
all.equal(ccf$value,
          vanillaOptionEuropean(S, X, tau, r, q, v, type = "call")$value)
all.equal(
  putCallParity("put", call=ccf$value, S=S, X=X, tau=tau, r=r, q=q),
  vanillaOptionEuropean(S, X, tau, r, q, v, type = "put")$value)
```

qTable

Prepare LaTeX Table with Quartile Plots

Description

The function returns the skeleton of a LaTeX tabular that contains the median, minimum and maximum of the columns of a matrix X. For each column, a quartile plot is added.

Usage

```
qTable(X, xmin = NULL, xmax = NULL, labels = NULL, at = NULL,
    unitlength = "5cm", linethickness = NULL,
    cnames = colnames(X), circlesize = 0.01,
    xoffset = 0, yoffset = 0, dec = 2, filename = NULL,
    funs = list(median = median, min = min, max = max),
    tabular.format, skip = TRUE)
```

qTable

Arguments

Х	a numeric matrix (or an object that can be coerced to a numeric matrix with as.matrix)
xmin	optional: the minimum for the x-axis. See Details.
xmax	optional: the maximum for the x-axis. See Details.
labels	optional: labels for the x-axis.
at	optional: where to put labels.
unitlength	the unitlength for LaTeX's picture environment. See Details.
linethickness	the linethickness for LaTeX's picture environment. See Details.
cnames	the column names of X
circlesize	the size of the circle in LaTeX's picture environment
xoffset	defaults to 0. See Details.
yoffset	defaults to 0. See Details.
dec	the number of decimals
filename	if provided, output is cat into a file
funs	A list of functions; the functions should be named. Default is
	list(median = median, min = min, max = max)
tabular.format	optional: character string like "rrrrr" that defines the format of the tabular.
skip	Adds a newline at the end of the tabular. Default is TRUE. (The behaviour prior to package version 0.27-0 corresponded to FALSE.)

Details

The function creates a one-column character matrix that can be put into a LaTeX file (the matrix holds a tabular). It relies on LaTeX's picture environment and should work for LaTeX and pdfLa-TeX. Note that the tabular needs generally be refined, depending on the settings and the data.

The tabular has one row for every column of X (and header and footer rows). A given row contains (per default) the median, the minimum and the maximum of the column; it also includes a picture environment the shows a quartile plot of the distribution of the elements in that column. Other functions can be specified via argument funs.

A number of parameters can be passed to LaTeX's picture environment: unitlength, xoffset, yoffset, linethickness. Sizes and lengths are functions of unitlength (linethickness is an exception; and while circlesize is a multiple of unitlength, it will not translate into an actual diameter of more than 14mm).

The whole tabular environment is put into curly brackets so that the settings do not change settings elsewhere in the LaTeX document.

If xmin, xmax, labels and at are not specified, they are computed through a call to pretty from the **base** package. If limits are specified, then both xmin and xmax must be set; if labels are used, then both labels and at must be specified.

To use the function in a vignette, use cat(tTable(X)) (and results=tex in the code chunk options). The vignette qTableEx shows some examples.

Value

A matrix of mode character. If filename is specified then qTable will have the side effect of writing a textfile with a LaTeX tabular.

Note

qTable returns a raw draft of a table for LaTeX. Please, spend some time on making it pretty.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Tufte, E. (2001) The Visual Display of Quantitative Information. 2nd edition, Graphics Press.

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

```
x <- rnorm(100, mean = 0, sd = 2)</pre>
y <- rnorm(100, mean = 1, sd = 2)
z <- rnorm(100, mean = 1, sd = 0.5)
X \leq cbind(x, y, z)
res <- qTable(X)</pre>
print(res)
cat(res)
## Not run:
## show vignette with examples
qt <- vignette("qTableEx", package = "NMOF")</pre>
print(qt)
edit(qt)
## create a simple LaTeX file 'test.tex':
## ---
## \documentclass{article}
## \begin{document}
##
     \input{res.tex}
## \end{document}
## ---
res <- qTable(X, filename = "res.tex", yoffset = -0.025, unitlength = "5cm",</pre>
              circlesize = 0.0125, xmin = -10, xmax = 10, dec = 2)
## End(Not run)
```

randomReturns

Description

Create a matrix of random returns.

Usage

randomReturns(na, ns, sd, mean = 0, rho = 0, exact = FALSE)

Arguments

na	number of assets
ns	number of return scenarios
sd	the standard deviation: either a single number or a vector of length na
mean	the mean return: either a single number or a vector of length na
rho	correlation: either a scalar (i.e. a constant pairwise correlation) or a correlation matrix
exact	logical: if TRUE, return a random matrix whose column means, standard de- viations and correlations match the specified values exactly (up to numerical precision)

Details

The function corresponds to the function random_returns, described in the second edition of NMOF (the book).

Value

a numeric matrix of size na times ns

Note

The function corresponds to the function random_returns, described in the second edition of NMOF (the book).

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

See Also

МC

Examples

repairMatrix Repair an Indefinite Correlation Matrix

Description

The function 'repairs' an indefinite correlation matrix by replacing its negative eigenvalues by zero.

Usage

repairMatrix(C, eps = 0)

Arguments

С	a correlation matrix
eps	a small number

Details

The function 'repairs' a correlation matrix: it replaces negative eigenvalues with eps and rescales the matrix such that all elements on the main diagonal become unity again.

Value

Returns a numeric matrix.

70

resampleC

Note

This function may help to cure a numerical problem, but it will rarely help to cure an empirical problem. (Garbage in, garbage out.)

See also the function nearPD in the Matrix package.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Rebonato, R. and Jaeckel, P. (1999) The most general methodology to create a valid correlation matrix for risk management and option pricing purposes.

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Examples

```
## example: build a portfolio of three assets
C <- c(1,.9,.9,.9,1,.2,.9,.2,1)
dim(C) <- c(3L, 3L)
eigen(C, only.values = TRUE)
vols <- c(.3, .3, .3)  ## volatilities
S <- C * outer(vols,vols)  ## covariance matrix
w <- c(-1, 1, 1)  ## a portfolio
w %*% S %*% w  ## variance of portfolio is negative!
sqrt(as.complex(w %*% S %*% w))
S <- repairMatrix(C) * outer(vols,vols)
w %*% S %*% w  ## more reasonable
sqrt(w %*% S %*% w)
```

resampleC

Resample with Specified Rank Correlation

Description

Resample with replacement from a number of vectors; the sample will have a specified rank correlation.

Usage

resampleC(..., size, cormat)

resampleC

Arguments

	numeric vectors; they need not have the same length.
size	an integer: the number of samples to draw
cormat	the rank correlation matrix

Details

See Gilli, Maringer and Schumann (2011), Section 7.1.2. The function samples with replacement from the vectors passed through The resulting samples will have an (approximate) rank correlation as specified in cormat.

The function uses the eigenvalue decomposition to generate the correlation; it will not break down in case of a semidefinite matrix. If an eigenvalue of cormat is smaller than zero, a warning is issued (but the function proceeds).

Value

a numeric matrix with size rows. The columns contain the samples; hence, there will be as many columns as vectors passed through

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

repairMatrix

```
## a sample
v1 <- rnorm(20)
v2 <- runif(50)
v3 <- rbinom(100, size = 50, prob = 0.4)
## a correlation matrix
cormat <- array(0.5, dim = c(3, 3))
diag(cormat) <- 1
cor(resampleC(a = v1, b = v2, v3, size = 100, cormat = cormat),
    method = "spearman")</pre>
```
restart0pt

Description

The function provides a simple wrapper for the optimisation algorithms in the package.

Usage

```
restartOpt(fun, n, OF, algo, ...,
    method = c("loop", "multicore", "snow"),
    mc.control = list(), cl = NULL,
    best.only = FALSE)
```

Arguments

fun	the optimisation function: DEopt, GAopt, LSopt, TAopt or PSopt
n	the number of restarts
OF	the objective function
algo	the list algo that is passed to the particular optimisation function
	additional data that is passed to the particular optimisation function
method	can be loop (the default), multicore or snow. See Details.
mc.control	a list containing settings that will be passed to mclapply if method is multicore. Must be a list of named elements. See the documentation of mclapply.
cl	default is NULL. If method snow is used, this must be a cluster object or an integer (the number of cores).
best.only	if TRUE, only the best run is reported. Default is FALSE.

Details

The function returns a list of lists. If a specific starting solution is passed, all runs will start from this solution. If this is not desired, initial solutions can be created randomly. This is done per default in DEopt, GAopt and PSopt, but LSopt and TAopt require to specify a starting solution.

In case of LSopt and TAopt, the passed initial solution algo\$x0 is checked with is.function: if TRUE, the function is evaluated in each single run. For DEopt, GAopt and PSopt, the initial solution (which also can be a function) is specified with algo\$initP.

The argument method determines how fun is evaluated. Default is loop. If method is "multicore", function mclapply from package **parallel** is used. Further settings for mclapply can be passed through the list mc.control. If multicore is chosen but the functionality is not available, then method will be set to loop and a warning is issued. If method == "snow", function clusterApply from package **parallel** is used. In this case, the argument cl must either be a cluster object (see the documentation of clusterApply) or an integer. If an integer, a cluster will be set up via makeCluster(c(rep("localhost", cl)), type = "SOCK"), and stopCluster is called when the function is exited. If snow is chosen but **parallel** is not available or cl is not specified, then method will be set to loop and a warning is issued. In case that cl is an cluster object, stopCluster will not be called automatically. 74

Value

If best.only is FALSE (the default), the function returns a list of n lists. Each of the n lists stores the output of one of the runs.

If best.only is TRUE, only the best restart is reported. The returned list has the structure specific to the used method.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

DEopt, GAopt, LSopt, PSopt, TAopt

Examples

```
## see example(DEopt)
algo <- list(nP = 50L,
              F = 0.5,
             CR = 0.9,
            \min = c(-10, -10),
            max = c(10, 10),
    printDetail = FALSE,
       printBar = FALSE)
## choose a larger 'n' when you can afford it
algo$nG <- 100L
res100 <- restartOpt(DEopt, n = 5L, OF = tfTrefethen, algo = algo)</pre>
res100F <- sapply(res100, `[[`, "OFvalue")</pre>
algo$nG <- 200L
res200 <- restartOpt(DEopt, n = 5L, OF = tfTrefethen, algo = algo)</pre>
res200F <- sapply(res200, `[[`, "OFvalue")</pre>
xx <- pretty(c(res100F, res200F, -3.31))</pre>
plot(ecdf(res100F), main = "optimum is -3.306",
     xlim = c(xx[1L], tail(xx, 1L)))
abline(v = -3.3069, col = "red") ## optimum
lines(ecdf(res200F), col = "blue")
legend(x = "right", box.lty = 0, , lty = 1,
      legend = c("optimum", "100 generations", "200 generations"),
      pch = c(NA, 19, 19), col = c("red", "black", "blue"))
```

Ritter

```
## a 'best-of-N' strategy: given a sample x of objective
## function values, compute the probability that, after N draws,
## we have at least one realisation not worse than X
x <- c(0.1,.3,.5,.5,.6)
bestofN <- function(x, N) {</pre>
    nx <- length(x)</pre>
    function(X)
        1 - (sum(x > X)/nx)^N
}
bestof2 <- bestofN(x, 2)
bestof5 <- bestofN(x, 5)</pre>
bestof2(0.15)
bestof5(0.15)
## Not run:
## with R >= 2.13.0 and the compiler package
algo$nG <- 100L
system.time(res100 <- restartOpt(DEopt, n = 10L, OF = tfTrefethen, algo = algo))</pre>
require("compiler")
enableJIT(3)
system.time(res100 <- restartOpt(DEopt, n = 10L, OF = tfTrefethen, algo = algo))</pre>
## End(Not run)
```

Ritter

Download Jay Ritter's IPO Data

Description

Download IPO data provided by Jay Ritter and transform them into a data frame.

Usage

Arguments

dest.dir	character: a path to a directory
url	the data URL

Details

The function downloads IPO data provided by Jay R. Ritter https://site.warrington.ufl.edu/ ritter/. Since the data are provided in Excel format, package **openxlsx** is required.

The downloaded Excel gets a date prefix (today in format YYYYMMDD) and is stored in directory dest.dir. Before any download is attempted, the function checks whether a file with today's prefix exist in dest.dir; if yes, this file is used.

Ritter

Value

a data.frame:

CUSIP	CUSIP
Offer date	a Date
Company name	character: Company name
Ticker	character: Ticker
Founding	Founding year
PERM	PERM
VC dummy	VC Dummy
Rollup	Rollup
Dual	Dual
Post-issue shares	
	Post-issue shares
Internet	Internet

Author(s)

Enrico Schumann

References

https://site.warrington.ufl.edu/ritter/ipo-data/

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

See Also

French, Shiller

Examples

```
## Not run:
archive.dir <- "~/Downloads/Ritter"
if (!dir.exists(archive.dir))
        dir.create(archive.dir)
Ritter(archive.dir)
```

End(Not run)

SA.info

Description

The function can be called from the objective and neighbourhood function during a run of SAopt; it provides information such as the current iteration, the current solution, etc.

Usage

SA.info(n = 0L)

Arguments

n

generational offset; see Details.

Details

This function is still experimental.

The function can be called in the neighbourhood function or the objective function during a run of SAopt. It evaluates to a list with information about the state of the optimisation run, such as the current iteration or the currently best solution.

SA. info relies on parent.frame to retrieve its information. If the function is called within another function within the neighbourhood or objective function, the argument n needs to be increased.

Value

A list

calibration	logical: whether the algorithm is calibrating the acceptance probability
iteration	current iteration
step	current step for the given temperature level
temperature	current temperature (the number, not the value)
xbest	the best solution found so far

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (**NMOF** Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

SAopt

See Also

SAopt, TA.info

Examples

MINIMAL EXAMPLE for SAopt

```
## the objective function evaluates to a constant
fun <- function(x)</pre>
    0
## the neighbourhood function does not even change
## the solution; it only reports information
nb <- function(x) {</pre>
    info <- SA.info()</pre>
    cat("current step ",
                                info$step,
        "| current iteration ", info$iteration, "\n")
    х
}
## run SA
algo <- list(nS = 5, nT = 2, nD = 10,
             initT = 1,
             x0 = rep(0, 5),
             neighbour = nb,
             printBar = FALSE)
ignore <- SAopt(fun, algo)</pre>
```

SAopt

Optimisation with Simulated Annealing

Description

The function implements a Simulated-Annealing algorithm.

Usage

SAopt(OF, algo = list(), ...)

Arguments

OF	The objective function, to be minimised. Its first argument needs to be a solution x; it will be called as $OF(x,)$.
algo	A list of settings for the algorithm. See Details.
	other variables passed to OF and algo\$neighbour. See Details.

SAopt

Details

Simulated Annealing (SA) changes an initial solution iteratively; the algorithm stops after a fixed number of iterations. Conceptually, SA consists of a loop than runs for a number of iterations. In each iteration, a current solution xc is changed through a function algo\$neighbour. If this new (or neighbour) solution xn is not worse than xc, ie, if $OF(xn, ...) \leq OF(xc, ...)$, then xn replaces xc. If xn is worse, it still replaces xc, but only with a certain probability. This probability is a function of the degree of the deterioration (the greater, the less likely the new solution is accepted) and the current iteration (the longer the algorithm has already run, the less likely the new solution is accepted).

The list algo contains the following items.

- nS The number of steps per temperature. The default is 1000; but this setting depends very much on the problem.
- nT The number of temperatures. Default is 10.
- nI Total number of iterations, with default NULL. If specified, it will override nS with ceiling(nI/nT). Using this option makes it easier to compare and switch between functions LSopt, TAopt and SAopt.
- nD The number of random steps to calibrate the temperature. Defaults to 2000.
- initT Initial temperature. Defaults to NULL, in which case it is automatically chosen so that initProb is achieved.
- finalT Final temperature. Defaults to 0.
- alpha The cooling constant. The current temperature is multiplied by this value. Default is 0.9.
- mStep Step multiplier. The default is 1, which implies constant number of steps per temperature. If greater than 1, the step number nS is increased to m*nS (and rounded).
- x0 The initial solution. If this is a function, it will be called once without arguments to compute an initial solution, ie, x0 <- algo\$x0(). This can be useful when the routine is called in a loop of restarts, and each restart is to have its own starting value.
- neighbour The neighbourhood function, called as neighbour(x, ...). Its first argument must be a solution x; it must return a changed solution.
- printDetail If TRUE (the default), information is printed. If an integer i greater then one, information is printed at very ith iteration.
- printBar If TRUE (default is FALSE), a txtProgressBar (from package utils) is printed. The progress bar is not shown if printDetail is an integer greater than 1.
- storeF if TRUE (the default), the objective function values for every solution in every generation are stored and returned as matrix Fmat.
- storeSolutions Default is FALSE. If TRUE, the solutions (ie, decision variables) in every generation are stored and returned in list xlist (see Value section below). To check, for instance, the current solution at the end of the ith generation, retrieve xlist[[c(2L, i)]].
- classify Logical; default is FALSE. If TRUE, the result will have a class attribute SAopt attached.
- OF.target Numeric; when specified, the algorithm will stop when an objective-function value as low as OF.target (or lower) is achieved. This is useful when an optimal objective-function value is known: the algorithm will then stop and not waste time searching for a better solution.

At the minimum, algo needs to contain an initial solution x0 and a neighbour function.

The total number of iterations equals algo\$nT times algo\$nS (plus possibly algo\$nD).

Value

SAopt returns a list with five components:

xbest	the solution
OFvalue	objective function value of the solution, ie, OF(xbest,)
Fmat	if algo\$storeF is TRUE, a matrix with one row for each iteration (excluding the initial algo\$nD steps) and two columns. The first column contains the objective function values of the neighbour solution at a given iteration; the second column contains the value of the current solution. Since SA can walk away from locally-optimal solutions, the best solution can be monitored through cummin(Fmat[,2L]).
xlist	if algo $storeSolutions$ is TRUE, a list; else NA. Contains the neighbour solutions at a given iteration (xn) and the current solutions (xc). Example: Fmat[i, 2L] is the objective function value associated with xlist[[c(2L, i)]].
initial.state	the value of .Random.seed when the function was called.

If algo\$classify was set to TRUE, the resulting list will have a class attribute TAopt.

Note

If the ... argument is used, then all the objects passed with ... need to go into the objective function and the neighbourhood function. It is recommended to collect all information in a list myList and then write OF and neighbour so that they are called as OF(x, myList) and neighbour(x, myList). Note that x need not be a vector but can be any data structure (eg, a matrix or a list).

Using an initial and final temperature of zero means that SA will be equivalent to a Local Search. The function LSopt may be preferred then because of smaller overhead.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Kirkpatrick, S., Gelatt, C.D. and Vecchi, M.P. (1983). Optimization with Simulated Annealing. Science. **220** (4598), 671–680.

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

LSopt, TAopt, restartOpt

SAopt

Examples

```
## Aim: given a matrix x with n rows and 2 columns,
        divide the rows of x into two subsets such that
##
##
        in one subset the columns are highly correlated,
##
        and in the other lowly (negatively) correlated.
        constraint: a single subset should have at least 40 rows
##
## create data with specified correlation
n <- 100L
rho <- 0.7
C <- matrix(rho, 2L, 2L); diag(C) <- 1
x <- matrix(rnorm(n * 2L), n, 2L) %*% chol(C)</pre>
## collect data
data <- list(x = x, n = n, nmin = 40L)
## a random initial solution
x0 <- runif(n) > 0.5
## a neighbourhood function
neighbour <- function(xc, data) {</pre>
   xn <- xc
   p <- sample.int(data$n, size = 1L)</pre>
   xn[p] \le abs(xn[p] - 1L)
    # reject infeasible solution
   c1 <- sum(xn) >= data$nmin
   c2 <- sum(xn) <= (data$n - data$nmin)
    if (c1 && c2) res <- xn else res <- xc
    as.logical(res)
}
## check (should be 1 FALSE and n-1 TRUE)
x0 == neighbour(x0, data)
## objective function
OF <- function(xc, data)
    -abs(cor(data$x[xc, ])[1L, 2L] - cor(data$x[!xc, ])[1L, 2L])
## check
OF(x0, data)
## check
OF(neighbour(x0, data), data)
## plot data
par(mfrow = c(1,3), bty = "n")
plot(data$x,
     xlim = c(-3,3), ylim = c(-3,3),
     main = "all data", col = "darkgreen")
## *Local Search*
algo <- list(nS = 3000L,
             neighbour = neighbour,
```

```
x0 = x0,
             printBar = FALSE)
sol1 <- LSopt(OF, algo = algo, data=data)</pre>
sol1$0Fvalue
## *Simulated Annealing*
algo$nT <- 10L
algo$nS <- ceiling(algo$nS/algo$nT)</pre>
sol <- SAopt(OF, algo = algo, data = data)</pre>
sol$OFvalue
c1 <- cor(data$x[ sol$xbest, ])[1L, 2L]</pre>
c2 <- cor(data$x[!sol$xbest, ])[1L, 2L]</pre>
lines(data$x[ sol$xbest, ], type = "p", col = "blue")
plot(data$x[ sol$xbest, ], col = "blue",
     xlim = c(-3, 3), ylim = c(-3, 3),
     main = paste("subset 1, corr.", format(c1, digits = 3)))
plot(data$x[!sol$xbest, ], col = "darkgreen",
     xlim = c(-3,3), ylim = c(-3,3),
     main = paste("subset 2, corr.", format(c2, digits = 3)))
## compare LS/SA
par(mfrow = c(1, 1), bty = "n")
plot(sol1$Fmat[ , 2L],type = "1", ylim=c(-1.5, 0.5),
     ylab = "OF", xlab = "Iterations")
lines(sol$Fmat[ , 2L],type = "1", col = "blue")
legend(x = "topright", legend = c("LS", "SA"),
       lty = 1, lwd = 2, col = c("black", "blue"))
```

```
Shiller
```

Download Robert Shiller's Data

Description

Download the data provided by Robert Shiller and transform them into a data frame.

Usage

Arguments

dest.dir	character: a path to a directory
url	the data URL

Shiller

Details

The function downloads US stock-market data provided by Robert Shiller which he used in his book 'Irrational Exhuberance'. Since the data are provided in Excel format, package **readxl** is required. The downloaded Excel gets a date prefix (today in format YYYYMMDD) and is stored in directory

dest.dir. Before any download is attempted, the function checks whether a file with today's prefix exist in dest.dir; if yes, the file is used.

Value

a data.frame:

Date	end of month
Price	numeric
Dividend	numeric
Earnings	numeric
CPI	numeric
Long Rate	numeric
Real Price	numeric
Real Dividend	numeric
Real Earnings	numeric
CAPE	numeric

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann.net/NMOF.htm#NMOFmanual

Shiller, R.J. (2015) Irrational Exhuberance. Princeton University Press. 3rd edition.

See Also

French

Examples

```
## Not run:
archive.dir <- "~/Downloads/Shiller"
if (!dir.exists(archive.dir))
        dir.create(archive.dir)
Shiller(archive.dir)
```

End(Not run)

showExample

Description

Display the code examples from 'Numerical Methods and Optimization and Finance'.

Usage

Arguments

file	a character vector of length one. See Details.
chapter	optional: a character vector of length one, giving the chapter name (see Details), or an integer, indicating a chapter number. Default is NULL: look in all chapters.
showfile	Should the file be displayed with file.show? Defaults to TRUE. A file will be displayed only if one single file only is identified by file and chapter.
includepaths	Should the file paths be displayed? Defaults to FALSE.
	Arguments passed to grep1; see Details.
edition	an integer: 1 and 2 are supported
search	a regular expression: search in the code files. Not supported yet.
ignore.case	passed to grep1; see Examples. Default is TRUE (which is much more helpful than the default FALSE before package version 2)

Details

showExample matches the specified file argument against the available file names via grepl(file, all.filenames, ignore.case = ignore.case, ...). If chapter is specified, a second match is performed, grepl(chapter, all.chapternames, ignore.case = ignore.case, ...). The chapternames are those in the book (e.g., 'Modeling dependencies'). The selected files are then those for which file name and chapter name could be matched.

Value

showExample returns a data.frame of at least two character vectors, Chapter and File. If includepaths is TRUE, Paths are included. If no file is found, the data.frame has zero rows. If a single file is identified and showfile is TRUE, the function has the side effect of displaying that file.

showChapterNames returns a character vector: the names of the book's chapters.

showExample

Note

The behaviour of the function changed slightly with version 2.0 to accommodate the code examples of the second edition of the book. Specifically, the function gained an argument edition, which defaults to 2. Also, the default for ignore.case was changed to TRUE. To get back the old behaviour of the function, set edition to 1 and ignore.case to FALSE.

The code files can also be downloaded from https://gitlab.com/NMOF.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2011) *Numerical Methods and Optimization in Finance*. Elsevier. doi:10.1016/C20090305693

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Examples

```
## list all files
showExample() ## 2nd edition is default
showExample(edition = 1)
## list specific files
showExample("Appendix")
showExample("Backtesting")
showExample("Heuristics")
showExample("tutorial") ## matches against filename
showExample(chapter = 13)
showExample(chapter = "tutorial")
## show where a file is installed
showExample(chapter = "portfolio", includepaths = TRUE)
## first edition
showExample("equations.R", edition = 1)
showExample("example", chapter = "portfolio", edition = 1)
showExample("example", chapter = 13, edition = 1)
showExample("example", chapter = showChapterNames(1)[13L], edition = 1)
```

TA.info

Description

The function can be called from the objective and neighbourhood function during a run of TAopt; it provides information such as the current iteration, the current solution, etc.

Usage

TA.info(n = 0L)

Arguments

n

generational offset; see Details.

Details

This function is still experimental.

The function can be called in the neighbourhood function or the objective function during a run of TAopt. It evaluates to a list with the state of the optimisation run, such as the current iteration.

TA. info relies on parent.frame to retrieve its information. If the function is called within another function in the neighbourhood or objective function, the argument n needs to be increased.

Value

A list

OF.sampling	logical: if TRUE, is the algorithm sampling the objective function to compute thresholds; otherwise (i.e. during the actual optimisation) FALSE
iteration	current iteration
step	current step (i.e. for a given threshold)
threshold	current threshold (the number, not the value)
xbest	the best solution found so far
OF.xbest	objective function value of best solution

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

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TAopt

See Also

TAopt

Examples

MINIMAL EXAMPLE for TAopt

```
## objective function evaluates to a constant
fun <- function(x)</pre>
   0
## neighbourhood function does not even change the solution,
## but it reports information
nb <- function(x) {</pre>
    tmp <- TA.info()</pre>
   cat("current threshold ",
                               tmp$threshold,
        "| current step ",
                               tmp$step,
        "| current iteration ", tmp$iteration, "\n")
    х
}
## run TA
algo <- list(nS = 5,
            nT = 2,
            nD = 3,
            x0 = rep(0, 5),
            neighbour = nb,
            printBar = FALSE,
            printDetail = FALSE)
ignore <- TAopt(fun, algo)</pre>
## printed output:
##
     current threshold NA | current step 1 | current iteration NA
##
      current threshold NA | current step 2 | current iteration NA
##
     current threshold NA | current step 3 | current iteration NA
     current threshold 1 | current step 1 | current iteration 1
##
##
     current threshold 1 | current step 2 | current iteration 2
##
     current threshold 1 | current step 3 | current iteration
                                                                   3
##
     current threshold 1 | current step 4 | current iteration
                                                                   4
     current threshold 1 | current step 5 | current iteration
##
                                                                   5
     current threshold 2 | current step 1 | current iteration
##
                                                                   6
##
     current threshold 2 | current step 2 | current iteration
                                                                   7
##
     current threshold
                        2 | current step 3 | current iteration
                                                                   8
##
      current threshold
                         2 | current step 4 | current iteration
                                                                   9
     current threshold 2 | current step 5 | current iteration 10
##
```

TAopt

Optimisation with Threshold Accepting

Description

The function implements the Threshold Accepting algorithm.

Usage

TAopt(OF, algo = list(), ...)

Arguments

OF	The objective function, to be minimised. Its first argument needs to be a solution x; it will be called as $OF(x,)$.
algo	A list of settings for the algorithm. See Details.
	other variables passed to OF and algo\$neighbour. See Details.

Details

Threshold Accepting (TA) changes an initial solution iteratively; the algorithm stops after a fixed number of iterations. Conceptually, TA consists of a loop than runs for a number of iterations. In each iteration, a current solution xc is changed through a function $algo\neighbour$. If this new (or neighbour) solution xn is not worse than xc, ie, if $OF(xn, ...) \leq OF(xc, ...)$, then xn replaces xc. If xn is worse, it still replaces xc as long as the difference in 'quality' between the two solutions is less than a threshold tau; more precisely, as long as $OF(xn, ...) - tau \leq OF(xc, ...)$. Thus, we also accept a new solution that is worse than its predecessor; just not too much worse. The threshold is typically decreased over the course of the optimisation. For zero thresholds TA becomes a stochastic local search.

The thresholds can be passed through the list algo (see below). Otherwise, they are automatically computed through the procedure described in Gilli et al. (2006). When the thresholds are created automatically, the final threshold is always zero.

The list algo contains the following items.

- nS The number of steps per threshold. The default is 1000; but this setting depends very much on the problem.
- nT The number of thresholds. Default is 10; ignored if algo\$vT is specified.
- nI Total number of iterations, with default NULL. If specified, it will override nS with ceiling(nI/nT). Using this option makes it easier to compare and switch between functions LSopt, TAopt and SAopt.
- nD The number of random steps to compute the threshold sequence. Defaults to 2000. Only used if algo\$vT is NULL.
- q The highest quantile for the threshold sequence. Defaults to 0.5. Only used if algo\$vT is NULL. If q is zero, TAopt will run with algo\$nT zero-thresholds (ie, like a Local Search).
- x0 The initial solution. If this is a function, it will be called once without arguments to compute an initial solution, ie, x0 <- algo\$x0(). This can be useful when the routine is called in a loop of restarts, and each restart is to have its own starting value.
- vT The thresholds. A numeric vector. If NULL (the default), TAopt will compute algo\$nT thresholds. Passing threshold can be useful when similar problems are handled. Then the time to sample the objective function to compute the thresholds can be saved (ie, we save algo\$nD

function evaluations). If the thresholds are computed and algo\$printDetail is TRUE, the time required to evaluate the objective function will be measured and an estimate for the remaining computing time is issued. This estimate is often very crude.

- neighbour The neighbourhood function, called as neighbour(x, ...). Its first argument must be a solution x; it must return a changed solution.
- printDetail If TRUE (the default), information is printed. If an integer i greater then one, information is printed at very ith iteration.
- printBar If TRUE (default is FALSE), a txtProgressBar (from package utils) is printed. The progress bar is not shown if printDetail is an integer greater than 1.
- scale The thresholds are multiplied by scale. Default is 1.
- drop0 When thresholds are computed, should zero values be dropped from the sample of objectivefunction values? Default is FALSE.
- stepUp Defaults to 0. If an integer greater than zero, then the thresholds are recycled, ie, vT is
 replaced by rep(vT, algo\$stepUp + 1) (and the number of thresholds will be increased by
 algo\$nT times algo\$stepUp). This option works for supplied as well as computed thresholds.
 Practically, this will have the same effect as restarting from a returned solution. (In Simulated
 Annealing, this strategy goes by the name of 'reheating'.)
- thresholds.only Defaults to FALSE. If TRUE, compute only threshold sequence, but do not actually run TA.
- storeF if TRUE (the default), the objective function values for every solution in every generation are stored and returned as matrix Fmat.
- storeSolutions Default is FALSE. If TRUE, the solutions (ie, decision variables) in every generation are stored and returned in list xlist (see Value section below). To check, for instance, the current solution at the end of the ith generation, retrieve xlist[[c(2L, i)]].
- classify Logical; default is FALSE. If TRUE, the result will have a class attribute TAopt attached. This feature is **experimental**: the supported methods (plot, summary) may change without warning.
- OF.target Numeric; when specified, the algorithm will stop when an objective-function value as low as OF.target (or lower) is achieved. This is useful when an optimal objective-function value is known: the algorithm will then stop and not waste time searching for a better solution.

At the minimum, algo needs to contain an initial solution x0 and a neighbour function.

The total number of iterations equals algo\$nT times (algo\$stepUp + 1) times algo\$nS (plus possibly algo\$nD).

Value

TAopt returns a list with four components:

xbest	the solution
OFvalue	objective function value of the solution, ie, OF(xbest,)
Fmat	if algo\$storeF is TRUE, a matrix with one row for each iteration (excluding the initial algo\$nD steps) and two columns. The first column contains the objective function values of the neighbour solution at a given iteration; the second column contains the value of the current solution. Since TA can walk away from locally-optimal solutions, the best solution can be monitored through cummin(Fmat[,2L]).

xlist	if algo\$storeSolutions is TRUE, a list; else NA. Contains the neighbour solu-
	tions at a given iteration (xn) and the current solutions (xc). Example: Fmat[i,
	2L] is the objective function value associated with xlist[[c(2L, i)]].
initial.state	the value of .Random.seed when the function was called.

If algo\$classify was set to TRUE, the resulting list will have a class attribute TAopt.

Note

If the ... argument is used, then all the objects passed with ... need to go into the objective function and the neighbourhood function. It is recommended to collect all information in a list myList and then write OF and neighbour so that they are called as OF(x, myList) and neighbour(x, myList). Note that x need not be a vector but can be any data structure (eg, a matrix or a list).

Using thresholds of size 0 makes TA run as a Local Search. The function LSopt may be preferred then because of smaller overhead.

Author(s)

Enrico Schumann

References

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See Also

LSopt, restartOpt. Simulated Annealing is implemented in function SAopt. Package **neighbours** (also on CRAN) offers helpers for creating neighbourhood functions.

TAopt

Examples

```
## Aim: given a matrix x with n rows and 2 columns,
        divide the rows of x into two subsets such that
##
##
        in one subset the columns are highly correlated,
##
        and in the other lowly (negatively) correlated.
        constraint: a single subset should have at least 40 rows
##
## create data with specified correlation
n <- 100L
rho <- 0.7
C <- matrix(rho, 2L, 2L); diag(C) <- 1
x <- matrix(rnorm(n * 2L), n, 2L) %*% chol(C)</pre>
## collect data
data <- list(x = x, n = n, nmin = 40L)
## a random initial solution
x0 <- runif(n) > 0.5
## a neighbourhood function
neighbour <- function(xc, data) {</pre>
   xn <- xc
   p <- sample.int(data$n, size = 1L)</pre>
   xn[p] \le abs(xn[p] - 1L)
    # reject infeasible solution
   c1 <- sum(xn) >= data$nmin
   c2 <- sum(xn) <= (data$n - data$nmin)
    if (c1 && c2) res <- xn else res <- xc
    as.logical(res)
}
## check (should be 1 FALSE and n-1 TRUE)
x0 == neighbour(x0, data)
## objective function
OF <- function(xc, data)
    -abs(cor(data$x[xc, ])[1L, 2L] - cor(data$x[!xc, ])[1L, 2L])
## check
OF(x0, data)
## check
OF(neighbour(x0, data), data)
## plot data
par(mfrow = c(1,3), bty = "n")
plot(data$x,
     xlim = c(-3,3), ylim = c(-3,3),
     main = "all data", col = "darkgreen")
## *Local Search*
algo <- list(nS = 3000L,
             neighbour = neighbour,
```

```
x0 = x0,
             printBar = FALSE)
sol1 <- LSopt(OF, algo = algo, data=data)</pre>
sol1$0Fvalue
## *Threshold Accepting*
algo$nT <- 10L
algo$nS <- ceiling(algo$nS/algo$nT)</pre>
sol <- TAopt(OF, algo = algo, data = data)</pre>
sol$OFvalue
c1 <- cor(data$x[ sol$xbest, ])[1L, 2L]</pre>
c2 <- cor(data$x[!sol$xbest, ])[1L, 2L]</pre>
lines(data$x[ sol$xbest, ], type = "p", col = "blue")
plot(data$x[ sol$xbest, ], col = "blue",
     xlim = c(-3,3), ylim = c(-3,3),
     main = paste("subset 1, corr.", format(c1, digits = 3)))
plot(data$x[!sol$xbest, ], col = "darkgreen",
     xlim = c(-3,3), ylim = c(-3,3),
     main = paste("subset 2, corr.", format(c2, digits = 3)))
## compare LS/TA
par(mfrow = c(1,1), bty = "n")
plot(sol1$Fmat[ ,2L],type="1", ylim=c(-1.5,0.5),
     ylab = "OF", xlab = "iterations")
lines(sol$Fmat[ ,2L],type = "1", col = "blue")
legend(x = "topright",legend = c("LS", "TA"),
       lty = 1, lwd = 2,col = c("black", "blue"))
```

testFunctions

Classical Test Functions for Unconstrained Optimisation

Description

A number of functions that have been suggested in the literature as benchmarks for unconstrained optimisation.

Usage

```
tfAckley(x)
tfEggholder(x)
tfGriewank(x)
tfRastrigin(x)
tfRosenbrock(x)
tfSchwefel(x)
tfTrefethen(x)
```

testFunctions

Arguments

х

a numeric vector of arguments. See Details.

Details

All functions take as argument only one variable, a numeric vector x whose length determines the dimensionality of the problem.

The Ackley function is implemented as

$$\exp(1) + 20 - 20 \exp\left(-0.2\sqrt{\frac{1}{n}\sum_{i=1}^{n}x_i^2}\right) - \exp\left(\frac{1}{n}\sum_{i=1}^{n}\cos(2\pi x_i)\right).$$

The minimum function value is zero; reached at x = 0.

The Eggholder takes a two-dimensional x, here written as x and y. It is defined as

$$-(y+47)\sin\left(\sqrt{|y+\frac{x}{2}+47|}\right) - x\sin\left(\sqrt{|x-(y+47)|}\right)$$
.

The minimum function value is -959.6407; reached at c(512, 404.2319).

The Griewank function is given by

$$1 + \frac{1}{4000} \sum_{i=1}^{n} x_i^2 - \prod_{i=1}^{n} \cos\left(\frac{x_i}{\sqrt{i}}\right) \,.$$

The function is minimised at x = 0; its minimum value is zero.

The *Rastrigin* function:

$$10n + \sum_{i=1}^{n} \left(x_i^2 - 10 \cos(2\pi x_i) \right) \,.$$

The minimum function value is zero; reached at x = 0. The *Rosenbrock* (or banana) function:

$$\sum_{i=1}^{n-1} \left(100(x_{i+1} - x_i^2)^2 + (1 - x_i)^2 \right) \,.$$

The minimum function value is zero; reached at x = 1. The *Schwefel* function:

$$\sum_{i=1}^n \left(-x_i \sin\left(\sqrt{|x_i|}\right) \right) \,.$$

The minimum function value (to about 8 digits) is -418.9829n; reached at x = 420.9687. *Trefethen*'s function takes a two-dimensional x (here written as x and y); it is defined as

$$\exp(\sin(50x)) + \sin(60e^y) + \sin(70\sin(x)) + \sin(\sin(80y)) - \sin(10(x+y)) + \frac{1}{4}(x^2+y^2)$$

The minimum function value is -3.3069; reached at c(-0.0244, 0.2106).

Value

The objective function evaluated at x (a numeric vector of length one).

Warning

These test functions represent *artificial* problems. It is practically not too helpful to fine-tune a method on such functions. (That would be like memorising all the answers to a particular multiple-choice test.) The functions' main purpose is checking the numerical implementation of algorithms.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

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See Also

DEopt, PSopt

Examples

persp for two-dimensional x

```
## Ackley
n <- 100L; surf <- matrix(NA, n, n)</pre>
x1 \le seq(from = -2, to = 2, length.out = n)
for (i in 1:n)
    for (j in 1:n)
        surf[i, j] <- tfAckley(c(x1[i], x1[j]))</pre>
persp(x1, x1, -surf, phi = 30, theta = 30, expand = 0.5,
      col = "goldenrod1", shade = 0.2, ticktype = "detailed",
      xlab = "x1", ylab = "x2", zlab = "-f", main = "Ackley (-f)",
      border = NA)
## Trefethen
n <- 100L; surf <- matrix(NA, n, n)</pre>
x1 \le seq(from = -10, to = 10, length.out = n)
for (i in 1:n)
    for (j in 1:n)
        surf[i, j] <- tfTrefethen(c(x1[i], x1[j]))</pre>
persp(x1, x1, -surf, phi = 30, theta = 30, expand = 0.5,
      col = "goldenrod1", shade = 0.2, ticktype = "detailed",
      xlab = "x1", ylab = "x2", zlab = "-f", main = "Trefethen (-f)",
      border = NA)
```

Description

Computes a portfolio similar to a benchmark, e.g. for tracking the benchmark's performance or identifying factors.

Usage

Arguments

var	the covariance matrix: a numeric (real), symmetric matrix. The first asset is the benchmark.
R	a matrix of returns: each colums holds the returns of one asset; each rows holds the returns for one observation. The first asset is the benchmark.
wmin	numeric: a lower bound on weights. May also be a vector that holds specific bounds for each asset.
wmax	numeric: an upper bound on weights. May also be a vector that holds specific bounds for each asset.
method	character. Currently, "qp" and "ls" are supported.
objective	character. Currently, "variance" and "sum.of.squares" are supported.
ls.algo	a list of named elements, for settings for method '1s'; see Details

Details

With method "qp", the function uses solve.QP from package **quadprog**. Because of the algorithm that solve.QP uses, var has to be positive definite (i.e. must be of full rank).

With method "ls", the function uses LSopt. Settings can be passed via ls.algo, which corresponds to LSopt's argument algo. Default settings are 2000 iterations and printBar, printDetail set to FALSE.

R is needed only when objective is "sum.of.squares" or method is 'ls'. (See Examples.)

Value

a numeric vector (the portfolio weights)

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

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See Also

minvar

Examples

```
if (requireNamespace("quadprog")) {
    ns <- 120
   R <- randomReturns(na = 1 + 20,
                        ns = ns,
                        sd = 0.03,
                        mean = 0.005,
                        rho = 0.7)
    var <- cov(R)</pre>
    sol.qp <- trackingPortfolio(var, wmax = 0.4)</pre>
    sol.ls <- trackingPortfolio(var = var, R = R, wmax = 0.4, method = "ls")</pre>
    data.frame(QP = round(100*sol.qp, 1),
               LS = round(100*sol.ls, 1))
    sol.qp <- trackingPortfolio(var, R = R, wmax = 0.4,</pre>
                                 objective = "sum.of.squares")
    sol.ls <- trackingPortfolio(var = var, R = R, wmax = 0.4, method = "ls",</pre>
                                 objective = "sum.of.squares")
    data.frame(QP = round(100*sol.qp, 1),
               LS = round(100*sol.ls, 1))
    ## same as 'sol.qp' above
    sol.qp.R <- trackingPortfolio(R = R,</pre>
                                      wmax = 0.4,
                                      objective = "sum.of.squares")
    sol.qp.var <- trackingPortfolio(var = crossprod(R),</pre>
                                      wmax = 0.4,
                                      objective = "variance")
    ## ==> should be the same
    all.equal(sol.qp.R, sol.qp.var)
}
```

vanillaBond

Description

Calculate the theoretical price and yield-to-maturity of a list of cashflows.

Usage

```
vanillaBond(cf, times, df, yields)
ytm(cf, times, y0 = 0.05, tol = 1e-05, maxit = 1000L, offset = 0)
duration(cf, times, yield, modified = TRUE, raw = FALSE)
convexity(cf, times, yield, raw = FALSE)
```

Arguments

cf	Cashflows; a numeric vector or a matrix. If a matrix, cashflows should be arranged in rows; times-to-payment correspond to columns.
times	times-to-payment; a numeric vector
df	discount factors; a numeric vector
yields	optional (instead of discount factors); zero yields to compute discount factor; if of length one, a flat zero curve is assumed
yield	numeric vector of length one (both duration and convexity assume a flat yield curve)
уØ	starting value
tol	tolerance
maxit	maximum number of iterations
offset	numeric: a 'base' rate over which to compute the yield to maturity. See Details and Examples.
modified	logical: return modified duration? (default TRUE)
raw	logical: default FALSE. Compute duration/convexity as derivative of cashflows' present value? Use this if you want to approximate the change in the bond price by a Taylor series (see Examples).

Details

vanillaBond computes the present value of a vector of cashflows; it may thus be used to evaluate not just bonds but any instrument that can be reduced to a deterministic set of cashflows.

ytm uses Newton's method to compute the yield-to-maturity of a bond (a.k.a. internal interest rate). When used with a bond, the initial outlay (i.e. the bonds dirty price) needs be included in the vector of cashflows. For a coupon bond, a good starting value y0 is the coupon divided by the dirty price of the bond.

An offset can be specified either as a single number or as a vector of zero rates. See Examples.

Value

numeric

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

See Also

NS, NSS

Examples

```
## ytm
cf <- c(5, 5, 5, 5, 5, 105) ## cashflows
times <- 1:6
                              ## maturities
y <- 0.0127
                             ## the "true" yield
b0 <- vanillaBond(cf, times, yields = y)</pre>
cf <- c(-b0, cf); times <- c(0, times)
ytm(cf, times)
## ... with offset
cf <- c(5, 5, 5, 5, 5, 105) ## cashflows
times <- 1:6
                              ## maturities
y <- 0.02 + 0.01
                             ## risk-free 2% + risk-premium 1%
b0 <- vanillaBond(cf, times, yields = y)</pre>
cf <- c(-b0, cf); times <- c(0, times)
ytm(cf, times, offset = 0.02) ## ... only the risk-premium
cf <- c(5, 5, 5, 5, 5, 105) ## cashflows
times <- 1:6
                              ## maturities
y <- NS(c(6,9,10,5)/100, times) ## risk-premium 1%</pre>
b0 <- vanillaBond(cf, times, yields = y + 0.01)</pre>
cf <- c(-b0, cf); times <- c(0, times)
ytm(cf, times, offset = c(0,y)) ## ... only the risk-premium
## bonds
cf <- c(5, 5, 5, 5, 105) ## cashflows
times <- 1:6
                             ## maturities
df <- 1/(1+y)^times
                             ## discount factors
all.equal(vanillaBond(cf, times, df),
          vanillaBond(cf, times, yields = y))
## ... using Nelson--Siegel
vanillaBond(cf, times, yields = NS(c(0.03,0,0,1), times))
```

several bonds

```
##
    cashflows are numeric vectors in a list 'cf',
     times-to-payment are are numeric vectors in a
##
##
    list 'times'
times <- list(1:3,</pre>
              1:4,
              0.5 + 0:5)
cf <- list(c(6, 6,
                             106),
           c(4, 4, 4,
                             104),
           c(2, 2, 2, 2, 2, 102))
alltimes <- sort(unique(unlist(times)))</pre>
M <- array(0, dim = c(length(cf), length(alltimes)))</pre>
for (i in seq_along(times))
    M[i, match(times[[i]], alltimes)] <- cf[[i]]</pre>
rownames(M) <- paste("bond.", 1:3, sep = "")</pre>
colnames(M) <- format(alltimes, nsmall = 1)</pre>
vanillaBond(cf = M, times = alltimes, yields = 0.02)
## duration/convexity
cf <- c(5, 5, 5, 5, 105) ## cashflows
times <- 1:6
                               ## maturities
                               ## yield to maturity
y <- 0.0527
d <- 0.001
                              ## change in yield (+10 bp)
vanillaBond(cf, times, yields = y + d) - vanillaBond(cf, times, yields = y)
duration(cf, times, yield = y, raw = TRUE) * d
duration(cf, times, yield = y, raw = TRUE) * d +
    convexity(cf, times, yield = y, raw = TRUE)/2 * d^2
```

vanillaOptionEuropean Pricing Plain-Vanilla (European and American) and Barrier Options (European)

Description

Functions to calculate the theoretical prices and (some) Greeks for plain-vanilla and barrier options.

Usage

> greeks = FALSE, model = NULL, ...)

Arguments

S	spot	
Х	strike	
Н	barrier	
tau	time-to-maturity in years	
r	risk-free rate	
q	continuous dividend yield, see Details.	
V	variance (volatility squared)	
tauD	vector of times-to-dividends in years. Only dividends with tauD greater than zero and not greater than tau are kept.	
D	vector of dividends (in currency units); default is no dividends.	
type	call or put; default is call.	
barrier.type	string: combination of up/down and in/out, such as downin	
rebate	currently not implemented	
greeks	compute Greeks? Defaults to TRUE. But see Details for American options.	
model	what model to use to value the option. Default is NULL which is equivalent to bsm.	
	parameters passed to pricing model	
М	number of time steps in the tree	
exercise	european (default) or american	
price	numeric; the observed price to be recovered through choice of volatility.	
uniroot.contro	1	
	A list. If there are elements named interval, tol or maxiter, these are passed to uniroot. Any other elements of the list are ignored.	
uniroot.info	logical; default is FALSE. If TRUE, the function will return the information re- turned by uniroot. See paragraph Value below.	

Details

For European options the formula of Messrs Black, Scholes and Merton is used. It can be used for equities (set q equal to the dividend yield), futures (Black, 1976; set q equal to r), currencies (Garman and Kohlhagen, 1983; set q equal to the foreign risk-free rate). For future-style options (e.g. options on the German Bund future), set q and r equal to zero.

The Greeks are provided in their raw ('textbook') form with only one exception: Theta is made negative. For practical use, the other Greeks are also typically adjusted: Theta is often divided by 365 (or some other yearly day count); Vega and Rho are divided by 100 to give the sensitivity for one percentage-point move in volatility/the interest rate. Raw Gamma is not much use if not adjusted for the actual move in the underlier.

For European options the Greeks are computed through the respective analytic expressions. For American options only Delta, Gamma and Theta are computed because they can be directly obtained from the binomial tree; other Greeks need to be computed through a finite difference (see Examples).

For the European-type options, the function understands vectors of inputs, except for dividends. American options are priced via a Cox-Ross-Rubinstein tree; no vectorisation is implemented here.

The implied volatility is computed with uniroot from the **stats** package (the default search interval is c(0.00001, 2); it can be changed through uniroot.control).

Dividends (D) are modelled via the escrowed-dividend model.

Value

Returns the price (a numeric vector of length one) if greeks is FALSE, else returns a list.

Note

If greeks is TRUE, the function will return a list with named elements (value, delta and so on). Prior to version 0.26-3, the first element of this list was named price.

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Haug, E. (2007) The Complete Guide to Option Pricing Formulas. 2nd edition. McGraw-Hill.

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

See Also

EuropeanCall, callCF

Examples

```
S <- 100; X <- 100; tau <- 1; r <- 0.02; g <- 0.06; vol <- 0.3
unlist(vanillaOptionEuropean(S, X, tau, r, q, vol^2, type = "put"))
S <- 100; X <- 110; tau <- 1; r <- 0.1; q <- 0.06; vol <- 0.3; type <- "put"
unlist(vanillaOptionAmerican(S, X, tau, r, q, vol^2, type = type,
                              greeks = TRUE))
## compute rho for 1% move
h <- 0.01
(vanillaOptionAmerican(S, X, tau, r + h, q, vol^2,
    type = type, greeks = FALSE) -
 vanillaOptionAmerican(S, X, tau, r, q, vol^2,
    type = type, greeks = FALSE)) / (h*100)
## compute vega for 1% move
h <- 0.01
(vanillaOptionAmerican(S, X, tau, r, q,(vol + h)<sup>2</sup>,
    type = type, greeks = FALSE) -
 vanillaOptionAmerican(S, X, tau, r, q, vol^2,
    type = type, greeks = FALSE)) / (h*100)
S <- 100; X <- 100
tau <- 1; r <- 0.05; q <- 0.00
D \le c(1,2); tauD \le c(0.3,.6)
type <- "put"
v <- 0.245^2 ## variance, not volatility</pre>
p <- vanillaOptionEuropean(S = S, X = X, tau, r, q, v = v,</pre>
                            tauD = tauD, D = D, type = type, greeks = FALSE)
vanillaOptionImpliedVol(exercise = "european", price = p,
     S = S, X = X, tau = tau, r = r, q = q, tauD = tauD, D = D, type = type)
p <- vanillaOptionAmerican(S = S, X = X, tau, r, q, v = v,</pre>
     tauD = tauD, D = D, type = type, greeks = FALSE)
vanillaOptionImpliedVol(exercise = "american", price = p,
     S = S, X = X, tau = tau, r = r, q = q, tauD = tauD, D = D, type =
     type, uniroot.control = list(interval = c(0.01, 0.5)))
## compute implied q
S <- 100; X <- 100
tau <- 1; r <- 0.05; q <- 0.072
v <- 0.22^2 ## variance, not volatility</pre>
call <- vanillaOptionEuropean(S=S, X = X, tau=tau, r=r, q=q, v=v,</pre>
         type = "call", greeks = FALSE)
put <- vanillaOptionEuropean(S=S, X = X, tau=tau, r=r, q=q, v=v,</pre>
         type = "put", greeks = FALSE)
```

xtContractValue

```
# ... the simple way
-(log(call + X * exp(-tau*r) - put) - log(S)) / tau
# ... the complicated way :-)
volDiffCreate <- function(exercise, call, put, S, X, tau, r) {</pre>
    f <- function(q) {</pre>
        cc <- vanillaOptionImpliedVol(exercise = exercise, price = call,</pre>
            S = S, X = X, tau = tau, r = r, q = q, type = "call")
        pp <- vanillaOptionImpliedVol(exercise = exercise, price = put,</pre>
            S = S, X = X, tau = tau, r = r, q = q, type = "put")
        abs(cc - pp)
    }
    f
}
f <- volDiffCreate(exercise = "european",</pre>
                   call = call, put = put, S = S, X = X, tau = tau, r)
optimise(f,interval = c(0, 0.2))$minimum
##
S <- 100; X <- 100
tau <- 1; r <- 0.05; q <- 0.072
v <- 0.22^2 ## variance, not volatility
vol <- 0.22
vanillaOptionEuropean(S=S, X = X, tau=tau, r=r, q=q, v=v,
                                                               ## with variance
                      type = "call", greeks = FALSE)
vanillaOptionEuropean(S=S, X = X, tau=tau, r=r, q=q, vol=vol, ## with vol
                      type = "call", greeks = FALSE)
vanillaOptionEuropean(S=S, X = X, tau=tau, r=r, q=q, vol=vol, ## with vol
                      type = "call", greeks = FALSE, v = 0.2^2)
```

xtContractValue Contract Value of Australian Government Bond Future

Description

Compute the contract value of an Australian government-bond future from its quoted price.

Usage

```
xtContractValue(quoted.price, coupon, do.round = TRUE)
xtTickValue(quoted.price, coupon, do.round = TRUE)
```

Arguments

quoted.price	The price, as in 99.02.
coupon	numeric; should be 6, not 0.06
do.round	If TRUE, round as done by ASX clearing house.

Details

Australian government-bond futures, traded at the Australian Securities Exchange (ASX), are quoted as 100 - yield. The function computes the actual contract value from the quoted price.

xtTickValue computes the tick value via a central difference.

Value

A numeric vector.

Author(s)

Enrico Schumann

References

https://www.rba.gov.au/mkt-operations/resources/tech-notes/pricing-formulae.html

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Schumann, E. (2023) Financial Optimisation with R (NMOF Manual). https://enricoschumann. net/NMOF.htm#NMOFmanual

Examples

xwGauss

Integration of Gauss-type

Description

Compute nodes and weights for Gauss integration.

Usage

```
xwGauss(n, method = "legendre")
changeInterval(nodes, weights, oldmin, oldmax, newmin, newmax)
```

xwGauss

Arguments

n	number of nodes	
method	character. default is "legendre"; also possible are "laguerre" and "hermite"	
nodes	the nodes (a numeric vector)	
weights	the weights (a numeric vector)	
oldmin	the minimum of the interval (typically as tabulated)	
oldmax	the maximum of the interval (typically as tabulated)	
newmin	the desired minimum of the interval	
newmax	the desired maximum of the interval	

Details

xwGauss computes nodes and weights for integration for the interval -1 to 1. It uses the method of Golub and Welsch (1969).

changeInterval is a utility that transforms nodes and weights to an arbitrary interval.

Value

a list with two elements

weights	a numeric vector
nodes	a numeric vector

Author(s)

Enrico Schumann

References

Gilli, M., Maringer, D. and Schumann, E. (2019) *Numerical Methods and Optimization in Finance*. 2nd edition. Elsevier. doi:10.1016/C2017001621X

Golub, G.H. and Welsch, J.H. (1969). Calculation of Gauss Quadrature Rules. *Mathematics of Computation*, **23**(106), pp. 221–230+s1–s10.

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See Also

callHestoncf

Examples

```
## examples from Gilli/Maringer/Schumann (2019), ch. 17
## a test function
f1 <- function(x) exp(-x)
m <- 5; a <- 0; b <- 5
h <- (b - a)/m
## rectangular rule -- left
w <- h; k <- 0:(m-1); x <- a + k * h
sum(w * f1(x))
## rectangular rule -- right
w <- h; k <- 1:m ; x <- a + k * h
sum(w * f1(x))
## midpoint rule
w <- h; k <- 0:(m-1); x <- a + (k + 0.5)*h
sum(w * f1(x))
## trapezoidal rule
w <- h
k <- 1:(m-1)
x <- c(a, a + k*h, b)
aux <- w * f1(x)
sum(aux) - (aux[1] + aux[length(aux)])/2
## R's integrate (from package stats)
integrate(f1, lower = a,upper = b)
## Gauss--Legendre
temp <- xwGauss(m)</pre>
temp <- changeInterval(temp$nodes, temp$weights,</pre>
                       oldmin = -1, oldmax = 1, newmin = a, newmax = b)
x <- temp$nodes; w <- temp$weights</pre>
sum(w * f1(x))
```

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