

Package ‘SACOBRA’

August 28, 2019

Type Package

Title Self-Adjusting COBRA

Version 1.1

Date 2019-08-15

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Description Performs surrogate-assisted optimization for expensive black-box constrained problems.

License GPL (>= 2)

Depends R (>= 2.14.0),

Suggests nloptr, FNN, MASS, dfoptim, DEoptim, lhs, rgl, grDevices,
scales, numDeriv, pracma, reshape2, data.table

Imports testit, methods, mgcv, R6

Collate 'cobraInit.R' 'cobraPhaseI.R' 'cobraPhaseII.R' 'debugModel.R'
'defaultDebugRBF.R' 'defaultRI.R' 'defaultSAC.R' 'defaultTR.R'
'defaultCA.R' 'drawSurrogate3d.R' 'evalReal.R' 'fnArchive.R'
'getPredY.R' 'initialHjkb.R' 'innerFuncs.R' 'isres2.R'
'modifyEquCons.R' 'modelSelection.R' 'multiRunPlot.R'
'multiRunPlot_2.R' 'multiCOBRA.R' 'nmkb2.R' 'RbfInter.R'
'repairChootinan.R' 'repairInfeasRI2.R' 'SACOBRA.R'
'startCobra.R' 'trainSurrogates.R' 'trustRegion.R'
'updateSaveCobra.R' 'Gproblems.R'

RoxygenNote 6.0.1

NeedsCompilation no

Repository CRAN

Date/Publication 2019-08-27 23:50:02 UTC

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SACOBRA-package

Self-adjusting Constrained Optimization with RBF Surrogates

Description

Self-adjusting Constrained Optimization with RBF Surrogates

Details

Package: SACOBRA
 Type: Package
 Version: 1.1
 Date: 16.08.2019
 License: GPL (>= 2)
 LazyLoad: yes

SACOBRA is a package for numeric constrained optimization of expensive black-box functions under severely limited budgets. The problem to solve is:

$$\begin{aligned}
 &\text{Minimize} && f(\vec{x}), \vec{x} \in [\vec{a}, \vec{b}] \subset \mathbf{R}^d \\
 &\text{subject to} && g_i(\vec{x}) \leq 0, i = 1, \dots, m \\
 &&& h_j(\vec{x}) = 0, j = 1, \dots, r.
 \end{aligned}$$

SACOBRA is an extension of the COBRA algorithm by Regis (R. Regis: "Constrained optimization by radial basis function interpolation for high-dimensional expensive black-box problems with infeasible initial points", Engineering Optimization, Taylor & Francis, 46, p. 218-243, 2013)

These extensions include:

- 1) A repair algorithm for infeasible solutions,
 - 2) an algorithm for handling equality constraints,
 - 3) several internal optimizers and several initial design generation methods,
 - 4) self-adjusting random restart algorithm,
 - 5) self-adjusting logarithmic transform for objective functions with large output ranges,
 - 6) range normalization of constraint functions,
 - 7) self-adjusting DRC (distance requirement cycle) selection,
 - 8) online model selection to select the best type of RBF for objective and constraint functions,
 - 9) online whitening for unconstrained optimization of functions with high conditioning.
- (Please note that the online whitening implementation is still underway and at this stage it is not recommended to be applied to expensive problems)

SACOBRA performs optimization with a minimum of true function evaluations. It has proven to work well on problems with high dimensions (e.g. $d=124$) and many constraints (e.g. 60). It is usable for all kind of numerical optimization of continuous functions, but not for combinatorial optimization.

For more details see:

- Bagheri, S.; Konen, W.; Emmerich, M.; Baeck, T.: "Self-adjusting parameter control for surrogate-assisted constrained optimization under limited budgets". In: Journal of Applied Soft Computing, Band 61, pages 377-393, 2017, https://www.researchgate.net/publication/319012980_Self-adjusting_parameter_control_for_surrogate-assisted_constrained_optimization_under_limited_budgets
- Bagheri, S.; Konen, W.; Baeck, T.: "Online selection of surrogate models for constrained black-box optimization". In: IEEE Symposium Series on Computational Intelligence (SSCI), 2016, <http://www.gm.fh-koeln.de/~konen/Publikationen/Bagh16-SSCI.pdf>

- Koch, P.; Bagheri, S.; Konen, W. et al.: "A New Repair Method For Constrained Optimization". In: Proceedings of the 17th Genetic and Evolutionary Computation Conference, 2015, <http://www.gm.fh-koeln.de/~konen/Publikationen/Koch2015a-GECCO.pdf>
- Koch, P.; Bagheri, S. et al.: "Constrained Optimization with a Limited Number of Function Evaluations" In: W. Hoffmann, F. & Huellermeier, E. (Eds.), Proceedings 24. Workshop Computational Intelligence, Universitaetsverlag Karlsruhe, 2014, 119-134, <http://www.gm.fh-koeln.de/~konen/Publikationen/Koch2014a-GMA-CI.pdf>.

The main entry point functions are `cobraInit` and `startCobra`. See `cobraInit` for an overview of adjustable SACOBRA-parameters. Examples are found in

- `startCobra`: solve a 13d-problem with 9 inequality constraints (G01)
- `cobraInit`: a problem with equality constraint
- `cobraPhaseII`: unconstrained sphere problem
- `multiCOBRA`: solve G11 problem nrun=4 times
- `COP`: load and solve G24, load and solve the scalable problem G03 with d=3

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References

<http://lwibs01.gm.fh-koeln.de/blogs/ciop/research/monrep/>

cobraInit

Initial phase for SACOBRA optimizer

Description

In this phase the important parameters are set and the initial design population are evaluated on the real function. The problem to solve is:

$$\text{Minimize } f(\vec{x}), \vec{x} \in [\vec{a}, \vec{b}] \subset \mathbf{R}^d$$

$$\text{subject to } g_i(\vec{x}) \leq 0, i = 1, \dots, m$$

$$h_j(\vec{x}) = 0, j = 1, \dots, r.$$

Usage

```
cobraInit(xStart, fn, fName, lower, upper, feval, initDesign = "LHS",
  initDesPoints = 2 * length(xStart) + 1, initDesOptP = NULL,
  initBias = 0.005, skipPhaseI = TRUE, seqOptimizer = "COBYLA",
  seqFeval = 1000, seqTol = 1e-06, ptail = TRUE, squares = TRUE,
  conTol = 0, DOSAC = 1, sac = defaultSAC(DOSAC), repairInfeas = FALSE,
  ri = defaultRI(), RBFmodel = "cubic", RBFwidth = -1,
  GaussRule = "One", widthFactor = 1, RBFrho = 0, MS = defaultMS(),
  equHandle = defaultEquMu(), rescale = TRUE, newlower = -1,
  newupper = 1, XI = DRCL, TrustRegion = FALSE, TRlist = defaultTR(),
  conditioningAnalysis = defaultCA(), penaF = c(3, 1.7, 3e+05),
  sigmaD = c(3, 2, 100), constraintHandling = "DEFAULT", verbose = 1,
  verboseIter = 10, DEBUG_RBF = defaultDebugRBF(), DEBUG_TR = FALSE,
  DEBUG_TRU = FALSE, DEBUG_RS = FALSE, DEBUG_XI = FALSE,
  trueFuncForSurrogates = FALSE, saveIntermediate = FALSE,
  saveSurrogates = FALSE, epsilonInit = NULL, epsilonMax = NULL,
  solu = NULL, cobraSeed = 42)
```

Arguments

xStart	a vector of dimension d containing the starting point for the optimization problem
fn	objective and constraint functions: fn is a function accepting a d -dimensional vector \vec{x} and returning an $(1+m+r)$ -dimensional vector $c(f, g_1, \dots, g_m, h_1, \dots, h_r)$
fName	the results of cobraPhaseII are saved to <code><fname>.Rdata</code>
lower	lower bound \vec{a} of search space, same dimension as xStart
upper	upper bound \vec{b} of search space, same dimension as xStart
feval	maximum number of function evaluations
initDesign	["LHS"] one out of ["RANDOM", "LHS", "BIASED", "OPTIMIZED", "OPTBIASED"]
initDesPoints	[$2*d+1$] number of initial points, must be smaller than feval
initDesOptP	[NULL] only for <code>initDesign=="OPTBIASED"</code> : number of points for the "OPT" phase. If NULL, take <code>initDesPoints</code> .
initBias	[0.005] bias for normal distribution in "OPTBIASED" and "BIASED"
skipPhaseI	[TRUE] if TRUE, then skip cobraPhaseI
seqOptimizer	["COBYLA"] string defining the optimization method for COBRA phases I and II, one out of ["COBYLA", "ISRES", "HJKB", "NMKB", "ISRESCOBY"]
seqFeval	[1000] maximum number of function evaluations on the surrogate model
seqTol	[$1e-6$] convergence tolerance for sequential optimizer, see param <code>tol</code> in nmkb or param <code>control\$xtol_rel</code> in cobylna
ptail	[TRUE] TRUE: with, FALSE: without polynomial tail in <code>trainRBF</code>
squares	[TRUE] set to TRUE for including the second order polynomials in building the fitness and constraint surrogates in <code>trainRBF</code>
conTol	[0.0] constraint violation tolerance

DOSAC	[1] set one out of [0112]. 0: COBRA-R settings, 1: SACOBRA settings, 2: SACOBRA settings with fewer parameters. The precise settings are documented in <code>defaultSAC</code> .
sac	<code>[defaultSAC(DOSAC)]</code> list with other parameters for SACOBRA.
repairInfeas	[FALSE] if TRUE, trigger the repair of appropriate infeasible solutions
ri	<code>[defaultRI()]</code> list with other parameters for <code>repairInfeasRI2</code>
RBFmodel	["cubic"] a string for the type of the RBF model, "cubic", "Gaussian" or "MQ"
RBFwidth	[-1] only relevant for Gaussian RBF model. Determines the width σ . For more details see parameter width in <code>trainGaussRBF</code> in <code>RBFinter.R</code> .
GaussRule	["One"] only relevant for Gaussian RBF model, see <code>trainGaussRBF</code>
widthFactor	[1.0] only relevant for Gaussian RBF model. Additional constant factor applied to each width σ
RBFrho	[0.0] experimental: 0: interpolating, > 0, approximating (spline-like) Gaussian RBFs
MS	<code>[defaultMS()]</code> list of online model selection parameters described in <code>defaultMS</code> . If <code>MS\$active = TRUE</code> then the type of RBF models for each function will be selected automatically and the <code>RBFmodel</code> parameter becomes irrelevant.
equHandle	<code>[defaultEquMu()]</code> list with of parameters for equality constraint handling described in <code>defaultEquMu()</code> . <code>equHandle\$active</code> is set to TRUE by default.
rescale	[TRUE] if TRUE, transform the input space from <code>[lower, upper]</code> to hypercube <code>[newlower, newupper]^d</code>
newlower	[-1] lower bound of each rescaled input space dimension, if <code>rescale==TRUE</code>
newupper	[+1] upper bound of each rescaled input space dimension, if <code>rescale==TRUE</code>
XI	[DRCL] magic parameters for the distance requirement cycle (DRC)
TrustRegion	[FALSE] if TRUE, perform trust region algorithm <code>trustRegion</code> .
TRlist	<code>[defaultTR()]</code> a list of parameters, needed only in case <code>TrustRegion==TRUE</code> .
conditioningAnalysis	<code>[defaultCA()]</code> A list with setting for the objective function conditioning analysis and online whitening
penaF	<code>[c(3,1.7,3e5)]</code> parameters for dynamic penalty factor (fct subProb in <code>cobraPhaseII</code>): <code>c(start, augment, max)</code> , only relevant if <code>seqOptimizer==HJKB</code> or <code>seqOptimizer==NMKB</code>
sigmaD	<code>[c(3,2.0,100)]</code> parameters for dynamic distance factor (fct subProb in <code>cobraPhaseII</code>): <code>c(start, augment, max)</code> , only relevant if <code>seqOptimizer==HJKB</code> or <code>seqOptimizer==NMKB</code>
constraintHandling	["DEFAULT"] (other choices: "JOINESHOUCK", "SMITHTATE", "COIT", "BAECKKHURI"; experimental, only relevant if <code>seqOptimizer==HJKB</code> or <code>seqOptimizer==NMKB</code> see the code in function <code>subProb</code> in <code>cobraPhaseII</code>)
verbose	[1] set one out of [0112], how much output to print
verboseIter	[10] an integer value. Printing the summarized results after each <code>verboseIter</code> iterations.

DEBUG_RBF	[defaultDebugRBF()] list with settings for visualization RBF (only for d==2)
DEBUG_TR	[FALSE] prints information about trust region status and visualisation for d==2 (coming soon)
DEBUG_TRU	[FALSE] visualize trust-region RBF (only for dimension==2)
DEBUG_RS	[FALSE] prints the RS probability in each iteration in the console
DEBUG_XI	[FALSE] if TRUE, then print in cobraPhaseII extra debug information: xStart in every iteration to console and add some extra debug columns to cobra\$df
trueFuncForSurrogates	[FALSE] if TRUE, use the true (constraint & fitness) functions instead of surrogates (only for debug analysis)
saveIntermediate	[FALSE] if TRUE, then cobraPhaseII saves intermediate results in dir 'results/' (create it, if necessary)
saveSurrogates	[FALSE] if TRUE, then cobraPhaseII returns the last surrogate models in cobra\$fitnessSurrogate and cobra\$constraintSurrogates
epsilonInit	[NULL] initial constant added to each constraint to maintain a certain margin to boundary
epsilonMax	[NULL] maximum for constant added to each constraint
solu	[NULL] the best-known solution (only for diagnostics). This is normally a vector of length d. If there are multiple solutions, it is a matrix with d columns (each row is a solution). If NULL, then the current best point will be used in cobraPhaseII . solu is given in original input space.
cobraSeed	[42] seed for random number generator

Details

If epsilonInit or epsilonMax are NULL on input, then cobra\$epsilonInit and cobra\$epsilonMax, resp., are set to $0.005 \cdot l$ where l is the smallest side of the search box.

Note that the parameters penaF, sigmaD, constraintHandling are only relevant for penalty-based internal optimizers [nmkb](#) or [HJKB](#). They are NOT relevant for default optimizer [coby1a](#).

Although the software was originally designed to handle only constrained optimization problems, it can also address unconstrained optimization problems

How to code which constraint is equality constraint? - Function fn should return an $(1 + m + r)$ -dimensional vector with named elements. The first element is the objective, the other elements are the constraints. All equality constraints should carry the name equ. (Yes, it is possible that multiple elements of a vector have the same name.)

Value

cobra, an object of class COBRA, this is a (long) list containing most of the argument settings (see above) and in addition (among others):

A	(feval x dim)-matrix containing the initial design points in input . space. If rescale==TRUE, all points are in rescaled input space.
Fres	a vector of the objective values of the initial design points

Gres	a matrix of the constraint values of the initial design points
nConstraints	the total number $m + r$ of constraints
Tfeas	the threshold parameter for the number of consecutive iterations that yield feasible solutions before margin epsilon is reduced
Tinfeas	the threshold parameter for the number of consecutive iterations that yield infeasible solutions before margin epsilon is increased
numViol	number of constraint violations
maxViol	maximum constraint violation
trueMaxViol	maximum constraint violation
trustregX	A vector of all refined solutions generated by trust region algorithm (see trustRegion)

Note that `cobra$Fres`, `cobra$fbest`, `cobra$fbestArray` and similar contain always the objective values of the original function `cobra$fn[1]`. (The surrogate models may be trained on a [plug-transformed](#) version of this function.)

Author(s)

Wolfgang Konen, Samineh Bagheri, Patrick Koch, Cologne University of Applied Sciences

See Also

[startCobra](#), [cobraPhaseI](#), [cobraPhaseII](#)

Examples

```
## Initialize cobra. The problem to solve is the sphere function sum(x^2)
## with the equality constraint that the solution is on a circle with
## radius 2 and center at c(1,0).
d=2
fName="onCircle"
cobra <- cobraInit(xStart=rep(5,d), fName=fName,
                  fn=function(x){c(obj=sum(x^2),equ=(x[1]-1)^2+(x[2]-0)^2-4)},
                  lower=rep(-10,d), upper=rep(10,d), feval=40)

## Run sacobra optimizer
cobra <- cobraPhaseII(cobra)

## The true solution is at solu = c(-1,0) (the point on the circle closest
## to the origin) where the true optimum is fn(solu)[1] = optim = 1
## The solution found by SACOBRA:
print(getXbest(cobra))
print(getFbest(cobra))

## Plot the resulting error (best-so-far feasible optimizer result - true optimum)
## on a logarithmic scale:
optim = 1
plot(abs(cobra$df$Best-optim),log="y",type="l",ylab="error",xlab="iteration",main=fName)
```

cobraPhaseI *Find a feasible solution.*

Description

Find a feasible solution using the COBRA optimizer phase I by searching new infill points. Please note that this phase can be skipped by setting the `cobra$skipPhaseI` parameter to TRUE in the initialization phase `cobraInit()`

Usage

```
cobraPhaseI(cobra)
```

Arguments

`cobra` an object of class COBRA, this is a (long) list containing all settings from `cobraInit`

Value

`cobra`, an object of class COBRA

Author(s)

Wolfgang Konen, Samineh Bagheri, Patrick Koch, Cologne University of Applied Sciences

See Also

[cobraPhaseII](#), [cobraInit](#)

cobraPhaseII *Improve the feasible solution by searching new infill points*

Description

Improve the feasible solution using the SACOBRA optimizer phase II by searching new infill points with the help of RBF surrogate models. May be even called if no feasible solution is found yet, then phase II will try to find feasible solutions.

The problem to solve iteratively is:

$$\begin{aligned} & \text{Minimize} && f(\vec{x}), \vec{x} \in [\vec{a}, \vec{b}] \subset \mathbf{R}^d \\ & \text{subject to} && g_i(\vec{x}) \leq 0, i = 1, \dots, m \\ & && h_j(\vec{x}) = 0, j = 1, \dots, r. \end{aligned}$$

In this phase the main optimization steps are repeated in a loop as long as the budget is not exhausted. In every iteration the surrogate models are updated and an optimization on the surrogates is done in order to find a better feasible solution.

Usage

```
cobraPhaseII(cobra)
```

Arguments

`cobra` an object of class COBRA, this is a (long) list containing all settings from `cobraInit`

Value

`cobra`, an object of class COBRA from `cobraInit`, enhanced here by the following elements (among others):

<code>fn</code>	function accepting a d -dimensional vector \vec{x} and returning an $(1+m+r)$ -vector $c(f, g_1, \dots, g_m, h_1, \dots, h_r)$. This function may be a rescaled and plog-transformed version of the original <code>fn</code> passed into <code>cobraInit</code> . The original <code>fn</code> is stored in <code>cobra\$originalFn</code> .
<code>df</code>	data frame with summary of the optimization run (see below)
<code>df2</code>	data frame with additional summary information (see below)
<code>dftr</code>	data frame with additional summary information for TR (see below)
<code>A</code>	$(\text{feval} \times d)$ -matrix containing all evaluated points in input space. If <code>rescale==TRUE</code> , all points are in rescaled input space.
<code>Fres</code>	a vector of the objective values of all evaluated points
<code>Gres</code>	a $(\text{feval} \times m)$ -matrix of the constraint values of all evaluated points
<code>predC</code>	a $(\text{feval} \times m)$ -matrix with the prediction of <code>cobra\$constraintSurrogates</code> at all evaluated points
<code>fbest</code>	the best feasible objective value found
<code>xbest</code>	the point in input space yielding the best feasible objective value
<code>ibest</code>	the corresponding iteration number (row of <code>cobra\$df</code> , of <code>cobra\$A</code>)
<code>PLOG</code>	If TRUE, then the objective surrogate model is trained on the plog -transformed objective function.

Note that `cobra$Fres`, `cobra$fbest`, `cobra$fbestArray` and similar contain always the objective values of the original function `cobra$fn[1]`. (The surrogate models may be trained on a **plog**-transformed version of this function.)

`feval = cobra$feval` is the maximum number of function evaluations.

The data frame `cobra$df` contains one row per iteration with columns

iter iteration index

y true objective value `Fres`

predY surrogate objective value. Note: The surrogate may be trained on plog-transformed training data, but `predY` is transformed back to the original objective range. NA for the initial design points.

predSolu surrogate objective value at best-known solution cobra\$solu, if given. If cobra\$solu is NULL, take the current point instead. Note: The surrogate may be trained on plog-transformed training data, but predSolu is transformed back to the original objective range. NA for the initial design points.

feasible boolean indicating the feasibility of infill point

feasPred boolean indicating if each infill point is feasible for cobra\$constraintSurrogates

nViolations number of violated constraints

maxViolation maximum constraint violation.

FEval number of function evaluations in sequential optimizer. NA if it was a repair step

Best ever-best feasible objective value fbest. As long as there is no feasible point, take among those with minimum number of violated constraints the one with minimum Fres.

optimizer e.g. "COBYLA"

optimizationTime in sec

conv optimizer convergence code

dist distance of the current point (row of cobra\$A) to the true solution cobra\$solu in rescaled space. If there is more than one solution, take the one which has the minimum distance element (since this is the solution to which the current run converges).

distOrig same as dist, but in original space

XI the DRC element used in the current iteration

seed the used seed in every run

The data frame cobra\$df2 contains one row per phase-II-iteration with columns

iter iteration index

predY surrogate objective value. Note: The surrogate may be trained on plog-transformed training data, but predY is transformed back to the original objective range. NA for the initial design points.

predVal surrogate objective value + penalty

predSolu surrogate objective value at true solution (see cobra\$df\$predSolu)

predSoluPenal surrogate objective value + penalty at true solution (only diagnostics)

sigmaD the sigmaD element used in the current iteration (see [cobraInit](#))

penaF penalty factor used in the current iteration (see [cobraInit](#))

XI the DRC element used in the current iteration

EPS the current used margin for constraint function modeling (see epsilonInit in [cobraInit](#))

Author(s)

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See Also

[cobraPhaseI](#), [cobraInit](#)

Examples

```
## Initialize cobra. The problem to solve is the unconstrained sphere function sum(x^2).

## In version 1.1 and higher there is no need for defining a dummy
## constraint function for the unconstrained problems
d=2
fName="sphere"
cobra <- cobraInit(xStart=rep(5,d), fName=fName,
                  fn=function(x){c(obj=sum(x^2))},
                  lower=rep(-10,d), upper=rep(10,d), feval=40)

## Run cobra optimizer
cobra <- cobraPhaseII(cobra)

## The true solution is at solu = c(0,0)
## where the true optimum is fn(solu)[1] = optim = 0
## The solution found by SACOBRA:
print(getXbest(cobra))
print(getFbest(cobra))

## Plot the resulting error (best-so-far feasible optimizer result - true optimum)
## on a logarithmic scale:
optim = 0
plot(cobra$df$Best-optim,log="y",type="l",ylab="error",xlab="iteration",main=fName)
```

COP

Constraint Optimization Problem Benchmark (G Function Suite)

Description

COP is an object of class `R6ClassGenerator` which can be used to access G problems (aka G functions) implementations in R, by simply generating a new instance of COP for each G function `problem<-COP.new("problem")`. The COP instances have the following useful attributes:

- name : name of the problem given by the user
- dimension: dimension of the problem. For the scalable problems G02 and G03, the dimension should be given by users, otherwise it will be set automatically
- lower: lower boundary of the problem
- upper: upper boundary of the problem
- fn: the COP function which can be passed to SACOBRA. (see fn description in [cobraInit](#))
- nConstraints: number of constraints
- xStart: The suggested optimization starting point
- solu: the best known solution, (only for diagnostics purposes)
- info: information about the problem

G function suite is a set of 24 constrained optimization problems with various properties like dimension, number of equality/ inequality constraint, feasibility ratio, etc. Although these problems were introduced as a suite in a technical report at CEC 2006, many of them have been used by different authors earlier.

For more details see: Liang, J., Runarsson, T.P., Mezura-Montes, E., Clerc, M., Suganthan, P., Coello, C.C., Deb, K.: Problem definitions and evaluation criteria for the CEC 2006 special session on constrained real-parameter optimization. *Journal of Applied Mechanics* 41, 8 (2006), http://www.lania.mx/~emezura/util/files/tr_cec06.pdf

Usage

COP

Format

An object of class R6ClassGenerator of length 24.

Author(s)

Samineh Bagheri, Wolfgang Konen

Examples

```
##creating an instance for G24 problem
G24<-COP$new("G24")

##initializing SACOBRA
cobra <- cobraInit(xStart=G24$lower, fName=G24$name,
                  fn=G24$fn,
                  lower=G24$lower, upper=G24$upper, feval=25)

## Run sacobra optimizer
cobra <- cobraPhaseII(cobra)

## The true solution is at solu = G24$solu
## The solution found by SACOBRA:
print(getXbest(cobra))
print(getFbest(cobra))
plot(abs(cobra$df$Best-G24$fn(G24$solu)[1]),log="y",type="l",
      ylab="error",xlab="iteration",main=G24$name)

## creating an instance for G03 in 2-dimensional space
G03<-COP$new("G03",2)

## Initializing sacobra
cobra <- cobraInit(xStart=G03$lower, fn=G03$fn,
                  fName=G03$name, lower=G03$lower, upper=G03$upper, feval=40)
```

defaultCA

*Default settings for online whitening functionality***Description**

Sets default values for the online whitening functionality in order to handle function with high conditioning. With the call `setOpts(cobra$CA, defaultCA())` it is possible to extend a partial list `cobra$CA` to a list containing all CA-elements (the missing ones are taken from `defaultCA()`).

As RBF interpolations face severe difficulties to deliver reasonable models for functions with high conditioning, we try to transform the function with high conditioning $f(\vec{x})$ to a better conditioned one $g(\vec{x})$ which is easier to model.

$$g(\vec{x}) = f(\mathbf{M}(\vec{x} - \vec{x}_c))$$

A possible transformation matrix \mathbf{M} is the squared inverse of the Hessian matrix $\mathbf{H}^{-0.5}$, assuming that \mathbf{M} is chosen with the following assumption:

$$\frac{\partial^2 g(\vec{x})}{\partial \vec{x}^2} = \mathbf{I}$$

Usage

```
defaultCA()
```

Details

The current version is only relevant for unconstrained problems. At this stage it is not recommended to apply the online whitening to expensive optimization problems as it demands a large number of function evaluations. Every online whitening call demands $4d^2 + 4d$ function evaluations.

Value

CA, a list of the following elements:

<code>active</code>	Set to TRUE if an online whitening of the fitness function is desired
<code>HessianType</code>	["real"] You can choose if the Hessian matrix is evaluated on the real function or on the surrogate model ["real", "surrogate"]. Please note that the determination of Hessian matrix on the real function at each point costs $4*d^2+d$ real function evaluations
<code>ITERS</code>	[seq(10,500,10)], pass a vector of integers to this parameter then the Hessian matrix will be updated only in the given iterations, we recommend applying the online-whitening every 10 iterations after the $10*d$ initial iterations. seq($10*d$, maxIter, 10), where d is the dimensionality of the optimization problem. If set as the character "all" then the Hessian matrix will be updated in each iteration and whitening procedure will be repeated
<code>alpha</code>	[1] you can assign any real value to this parameter. Only values between 0 to 2 are suggested. This value is used in order to modify the transformation center <code>tCenter</code> as follows: <code>xbest+alpha*(grad)</code> , and <code>grad</code> is the direction of the last improvement

See Also[setOpts](#)

defaultDebugRBF	<i>Default settings for debug visualization RBF (only for d==2)</i>
-----------------	---

Description

Sets default values for debug visualization RBF of SACOBRA.

Usage

```
defaultDebugRBF()
```

Value

DEBUG_RBF a list of the following elements:

active	If set to TRUE then debugVisualizeRBF is called every DEBUG_RBF\$every iterations
overlayTrueZ	If set to TRUE overlay the true objective function
DO_SNAPSHOT	do rgl.snapshot every DEBUG_RBF\$every iteration and store it in <code>sprintf("images.d/%s-%03d.png", cobra\$name, npts)</code>
every	Frequency of calling the debugVisualizeRBF function

See Also[debugVisualizeRBF](#)

defaultEquMu	<i>Default settings for equality handling mechanism</i>
--------------	---

Description

Sets suitable defaults for the equality handling part of SACOBRA.

The EH technique transforms each equality constraint $h(\vec{x}) = 0$ into two inequality constraints $h(\vec{x}) - \mu < 0$ and $-h(\vec{x}) - \mu < 0$ with an adaptively decaying margin μ .

If refine parameter is set to TRUE, then a refine mechanism is applied to shift the best found solution within the equality margin μ toward the feasible subspace by minimizing the sum of squared constraint surrogates with a conjugate gradient method.

$$\text{Minimize } \sum_i (\max(0, g_i(x)))^2 + \sum_j (h_j(x))^2$$

Usage

```
defaultEquMu()
```

Details

With the call `setOpts(equHandle, defaultEquMu())` it is possible to extend a partial list `equHandle` list which is set by user to a list containing all `equHandle`-elements (the missing ones are taken from `defaultEquMu()`). These settings are used by `cobraInit` for initializing the equality margin μ and by the internal functions `updateCobraEqu` and `modifyMu`. The minimization step of refine mechanism is done by L-BFGS-B method in `optim` function from `stats` package.

Value

`equHandle`, a list with the following elements:

<code>active</code>	[TRUE] if set to TRUE the equality-handling (EH) technique is activated. The EH technique transforms each equality constraint $h(\vec{x}) = 0$ into two inequality constraints $h(\vec{x}) - \mu < 0$ and $-h(\vec{x}) - \mu < 0$ with an adaptively decaying margin μ .
<code>equEpsFinal</code>	[1e-07] lower bound for margin μ . <code>equEpsFinal</code> should be set to a small but non-zero value (larger than machine accuracy).
<code>initType</code>	["TAV"] the equality margin μ can be initialized with one of these approaches: ["TAV" "TMV" "EMV" "useGrange"] TAV : (Total Absolute Violation) takes the median of the sum of violations of the initial population. TMV : (Total Maximum Violation) takes the median of the maximum violation of the initial population EMV : takes the median of the maximum violation of equality constraints of the initial population useGrange : takes the average of the ranges of the equality constraint functions
<code>epsType</code>	["SAexpFunc"] type of the function used to modify margin μ during the optimization process can be one of ["SAexpFunc" "expFunc" "Zhang" "CONS"]. see <code>modifyMu</code> .
<code>dec</code>	[1.5] decay factor for margin μ . see <code>modifyMu</code>
<code>refine</code>	[TRUE] enables the refine mechanism f the equality handling mechanism.
<code>refineMaxit</code>	maximum number of iterations used in the refine step. Note that the refine step runs on the surrogate models and does not impose any extra real function evaluation.

See Also

[updateCobraEqu](#), [modifyMu](#)

 defaultMS

Default settings for the model-selection part of SACOBRA.

Description

Sets default values for the model-selection part `cobra$MS` of SACOBRA.

It is shown that different types of RBFs can deliver different qualities in modeling different functions. Using the online model selection functionality boosted the overall performance of SACOBRA on a large set of constrained problems. The algorithm trains every function (objective and constraints) with a given pool of models including different RBF types and width parameters. The type of model which performs the best in the last iterations `WinS` will be selected for each function. The quality of the models are determined by different measures of approximation error in each iteration

$$f(\vec{x}_{new}) - s(\vec{x}_{new})$$

Usage

`defaultMS()`

Details

With the call `setOpts(MS, defaultMS())` it is possible to extend a partial list `MS` to a list containing all `MS`-elements (the missing ones are taken from `defaultMS()`).

NOTE: Because of common crash observation, it is not recommended to include Gaussian model in the set of models especially for problems which require more than 100 function evaluations.

Value

`MS`, a list with the following elements

<code>active</code>	[F] If set to TRUE then <code>selectModel</code> calculates the best model for each constraint(s)/objective function
<code>models</code>	[c("cubic","MQ")] a set of model types that will be used to build the pool of models. Three types of RBF are implemented "cubic", "Gaussian" and "MQ" (multiquadric). Users can select one or combination of these models. Users can select a set of "width parameters" for "MQ" and "Gaussian" by setting <code>widths</code> parameter.
<code>widths</code>	[c(0.01,0.1,1,10)] a set of values for width parameter of RBF models. Only relevant if <code>models</code> include "Gaussian" or "MQ".
<code>freq</code>	[1] controls how often <code>selectModel</code> is called. In every <code>freq</code> iterations all the selected models are trained for all constraint/objective function(s)
<code>slidingW</code>	[T] when set to FALSE it uses the information taken from all the past iterations to assess the quality of the models. When set to TRUE, activates the sliding window functionality and it takes the information of the last <code>WinS</code> iterations (see <code>WinS</code>).

WinS	[1] size of the sliding window
quant	[3] 3: median, 2:0.25, 4:0.75. The measure used to compare the quality of the model in the last window.
apply	[T] if set to FALSE then the selected models are not used during the optimization. Only for debugging purposes.
considerXI	[F] If set to T then a subset of the approximation errors which are related to the current (DRC element) are considered to make the model selection decision

Author(s)

Samineh Bagheri

See Also

[setOpts](#)

defaultRI	<i>Default settings for repairInfeasRI2 and repairChootinan.</i>
-----------	--

Description

Sets suitable defaults for the repair-infeasible part of SACOBRA.

With the call `setOpts(myRI,defaultRI())` it is possible to extend a partial list myRI to a list containing all ri-elements (the missing ones are taken from defaultRI())

Usage

```
defaultRI(repairMargin = 0.01)
```

Arguments

repairMargin [1e-2] repair only solutions whose infeasibility is less than this margin

Details

The **infeasibility** of a solution is its maximum constraint violation (0 for a feasible solution).

Value

a list with the following elements:

RIMODE	[2] one out of {0,1,2,3 } with 0,1: deprecated older versions of RI2, 2: the recommended RI2-case, see repairInfeasRI2 , 3: Chootinan's method, see repairChootinan
eps1	[1e-4] include all constraints not eps1-feasible into the repair mechanism
eps2	[1e-4] selects the solution with the shortest shift among all random realizations which are eps2-feasible

q	[3.0] draw coefficients α_k from uniform distribution $U[0, q]$
mmax	[1000] draw mmax random realizations
repairMargin	repair only solutions whose infeasibility is less than this margin.
repairOnlyFresBetter	[FALSE] if TRUE, then repair only iterates with fitness < so-far-best-fitness + marFres
marFres	[0.0] only relevant if repairOnlyFresBetter==TRUE

A solution x is said to be **ϵ -feasible** for constraint function f , if

$$f(x) + \epsilon \leq 0$$

Author(s)

Wolfgang Konen, Cologne University of Applied Sciences

See Also

[repairInfeasRI2](#), [repairChootinan](#)

defaultSAC	<i>Default settings for the SACOBRA part of SACOBRA.</i>
------------	--

Description

Sets suitable defaults for the SACOBRA part of SACOBRA.

With the call `setOpts(mySAC, defaultSAC())` it is possible to extend a partial list mySAC to a list containing all sac-elements (the missing ones are taken from defaultSAC()).

Usage

```
defaultSAC(DOSAC = 1)
```

Arguments

DOSAC	[0 1 2] with default 1. 0: COBRA-R settings (turn off SACOBRA), 1: SACOBRA settings, 2: SACOBRA settings with fewer parameters and more online adjustments (aFF and aCF are done parameter free).
-------	--

Details

For backward compatibility, a logical DOSAC (deprecated) is mapped from FALSE to 0 and from TRUE to 1.

Value

a list with the following elements (the values in parantheses [|] are the values for DOSAC=[0 | 1 | 2]):

RS	flag for random start algorithm [FALSE TRUE TRUE]
RStype	type of the function to calculate probability to start the internal optimizer with a random starting point[NA "SIGMOID" "CONSTANT"] (see function RandomStart in SACOBRA.R)
RSmax	maximum probability of a random start when RStype=="SIGMOID" (see RandomStart in SACOBRA.R). If RStype=="CONSTANT" then random start is done with a constant probability determined from mean(c(RSmax,RSmin)) [NA 0.3 0.3]
RSmin	minimum probability of a random start when RStype=="SIGMOID" (see RandomStart in SACOBRA.R) [NA 0.05 0.05]
RSAUTO	If TRUE then in every iteration where the fraction of feasible points in the population is smaller than 0.05, the RS probability is set to 0.3. [FALSE FALSE TRUE]
aDRC	flag for automatic DRC adjustment [FALSE TRUE TRUE]
aFF	flag for automatic objective function transformation [FALSE TRUE TRUE]
aCF	flag for automatic constraint function transformation [FALSE TRUE TRUE]
TFRange	threshold, if FRange is larger than TFRange, then apply automatic objective function transformation (see plog). [Inf 1e+05 -1]
TGR	threshold, if GRatio is larger than TGR, then apply automatic constraint function transformation. GRatio is the ratio "largest GRange / smallest GRange" where GRange is the min-max range of a specific constraint. If TGR < 1, then the transformation is always performed. [Inf 1e+03 -1].
Cs	If Cs iterations in a row do not improve the ever-best feasible solution, then perform a random restart. [10 10 10]
adaptivePLOG	(experimental) flag for objective function transformation with plog , where the parameter pShift is adapted during iterations. [FALSE FALSE FALSE]
onlinePLOG	flag for online decision making wether use plog or not according to p-effect plog . [FALSE FALSE TRUE]
pEffectInit	Initial pEffect value when using onlinePLOG. If pEffectInit >= 2 then the initial model is built after plog transformation. [NA NA 2]

Author(s)

Samineh Bagheri, Cologne University of Applied Sciences

See Also

[cobraInit](#), [cobraPhaseII](#)

defaultTR	<i>Default settings for the trust-region part of COBRA.</i>
-----------	---

Description

Sets default values for the trust-region part cobra\$TRlist of SACOBRA.

With the call `setOpts(myTR,defaultTR())` it is possible to extend a partial list myTR to a list containing all TR-elements (the missing ones are taken from defaultTR()).

Usage

defaultTR()

Value

a list with the following elements

shape	["cube"] Shape of the trust region can be chosen between cube or a sphere [cube sphere]
radiMin	[0.01] A value between 0 and 1, minimum fraction of the width of the search space to be used as radius of the trust region
radiMax	[0.8] A value between 0 and 1, maximum fraction of the width of the search space to be used as radius of the trust region
radiInit	[0.1] Initial radius of trust region
center	[cobra\$xbest] Center of the trust region can be the current bwst solution or the new solution[xbest xnew]

See Also

[setOpts](#), [trustRegion](#)

distLine	<i>Euclidean distance of x to all xp</i>
----------	--

Description

Euclidean distance of x to a line of points xp

Usage

distLine(x, xp)

Arguments

`x` vector of dimension `d`

`xp` `n` points x_i of dimension `d` are arranged in $(n \times d)$ matrix `xp`. If `xp` is a vector, it is interpreted as $(n \times 1)$ matrix, i.e. `d=1`.

Details

`distLine` is up to 40x faster than using `dist` and taking only the first row or column of the distance matrix returned.

Value

vector of length `n`, the Euclidean distances

DRCL *Distance Requirement Cycle, long version*

Description

Distance Requirement Cycle, long version: `c(0.3,0.05, 0.001, 0.0005,0.0)`

Usage

DRCL

Format

An object of class `numeric` of length 5.

DRCS *Distance Requirement Cycle, short version*

Description

Distance Requirement Cycle, short version: `c(0.001,0.0)`

Usage

DRCS

Format

An object of class `numeric` of length 2.

evalReal	<i>Evaluate new iterate on real function(s)</i>
----------	---

Description

Helper for [cobraPhaseII](#): The new iterate xNew, which was found by optimization on the surrogate models, is evaluated on the real function cobra\$fn. In the case of equality constraints, evalReal does the additional refine step (see Details).

Usage

```
evalReal(cobra, ev1, xNew, fValue, feval, optimConv, optimTime, currentEps,
         fitnessSurrogate = cobra$fitnessSurrogate)
```

Arguments

cobra	an object of class COBRA, this is a (long) list containing all settings from cobraPhaseII
ev1	a list, initially empty, gradually filled by calls to evalReal
xNew	the new point, see cobraPhaseII
fValue	fitness value estimated for xNew
feval	function evaluations on surrogates needed by COBRA optimizer
optimConv	see cobraPhaseII
optimTime	see cobraPhaseII
currentEps	artificial current margin for the equality constraints: A point is said to be artificially feasible , if $h_j(x) - currentEps \leq 0, -h_j(x) - currentEps \leq 0$, for all equality constraints and if it is feasible in the inequality constraints.
fitnessSurrogate	[cobra\$fitnessSurrogate] see cobraPhaseII

Details

If cobra\$equHandle\$active==TRUE, then xNew is first **refined**: The artificially feasible solution xNew is replaced by a refined solution ev1\$xNew. ev1\$xNew is created by using optim to minimize the function

$$\sum_i \max(0, g_i(x)) + \sum_j h_j^2(x)$$

Ideally, the refined solution ev1\$xNew should be **on** the equality constraints (within machine accuracy), but there is no guarantee that optim reaches this desired result.

Value

ev1, a list with the following n-dim vectors (n = number of iterations, the last element is from the new iterate / point xNew):

predY	prediction of fitnessSurrogate at xNew
predVal	fvalue (fitness + penalty in case of NMKB et al.)
feval	function evaluations on surrogates needed by COBRA optimizer
optimizerConvergence	see cobraPhaseII
optimizationTime	see cobraPhaseII
predC	prediction of cobra\$constraintSurrogates at xNew
feas	TRUE, if xNew is feasible for the current constraints
feasPred	TRUE, if xNew is feasible for cobra\$constraintSurrogates

In addition, ev1 has these elements:

xNew	d-dim vector, the new point, refined in the case of equality handling
xNewEval	cobra\$fn(xNew), an (1+nConstraints)-dim vector (objective,constraints)
newNumViol	scalar, the number of constraint violations (above cobra\$conTol) on true constraints from xNewEval
newNumPred	scalar, the number of constraint violations (above cobra\$conTol) on constraint surrogates for xNew
newMaxViol	scalar, the maximum constraint violation (with currentEps subtracted) on true constraints from xNewEval
trueMaxViol	scalar, the maximum constraint violation (w/o currentEps subtracted) on true constraints from xNewEval

If cobra\$equHandle\$active==TRUE, then the last four values are for xNew after the refine step. In this case, the first three elements newNumViol, newNumPred, and newMaxViol refer to the artificially enlarged equality constraints, i.e.

$$h_j(x) - currentEps \leq 0, -h_j(x) - currentEps \leq 0,$$

and the true inequality constraints $\max(0, g_i(x))$. The last element trueMaxViol measures the maximum violation among the true equality constraints $|h_j(x)|$ and the true inequality constraints $\max(0, g_i(x))$.

See Also

[cobraPhaseII](#)

forwardRescale	<i>Forward Rescaling</i>
----------------	--------------------------

Description

Scale vector x in original space forward to rescaled space (usually $[-1, 1]^d$)

Usage

```
forwardRescale(x, cobra)
```

Arguments

x	a vector in the original input space
cobra	list from cobraInit , we need here originalL a vector with lower bounds in original input space originalU a vector with upper bounds in original input space newlower a number, the rescaled lower bound for all dimensions newupper a number, the rescaled upper bound for all dimensions

Value

z , scaled version of vector x

See Also

[inverseRescale](#)

getFbest	<i>Return best objective function value</i>
----------	---

Description

Return the original objective function value at the best feasible solution

Usage

```
getFbest(cobra)
```

Arguments

cobra	an object of class COBRA (see cobraInit)
-------	---

Details

Note: We cannot take the best function value via `cobra$fn`, because this may be modified by `plog()` or others)

Value

the original objective function value at the best feasible solution

See Also

[getXbest](#)

getXbest

Return best feasible solution in original space

Description

Return best feasible solution in original space

Usage

```
getXbest(cobra)
```

Arguments

`cobra` an object of class COBRA (see [cobraInit](#))

Value

the best feasible solution in original space

See Also

[getFbest](#)

intern.archive.env *Archiving Environment*

Description

intern.archive.env is an independent environment where every evaluated point and its evaluation by the real function are stored in ARCHIVE and ARCHIVEY. This archive stores different values to cobra\$A and cobra\$Fres often during debugging and visualisation cases where the real function is evaluated very often for debugging purposes.

Usage

```
intern.archive.env
```

Format

An object of class environment of length 0.

interpRBF *Apply the trained cubic, MQ or Gaussian RBF interpolation to new data for $d > 1$.*

Description

Apply the trained cubic, MQ or Gaussian RBF interpolation to new data for $d > 1$.

Usage

```
interpRBF(x, rbf.model)
```

Arguments

x vector holding a point of dimension d
rbf.model trained RBF model (or set of models), see [trainCubicRBF](#) or [trainGaussRBF](#)

Value

value $s(\vec{x})$ of the trained model at \vec{x}
- or -
vector $s_j(\vec{x})$ with values for all trained models $j = 1, \dots, m$ at \vec{x}

Author(s)

Wolfgang Konen (<wolfgang.konen@th-koeln.de>)

See Also

[trainCubicRBF](#), [trainMQRBF](#), [trainGaussRBF](#), [predict.RBFinter](#)

inverseRescale *Inverse Rescaling*

Description

Scale vector x in rescaled space back to original space

Usage

```
inverseRescale(x, cobra)
```

Arguments

x a vector in the rescaled input space (usually $[-1, 1]^d$)

$cobra$ list from `cobraInit`, we need here

originalL a vector with lower bounds in original input space

originalU a vector with upper bounds in original input space

newlower a number, the rescaled lower bound for all dimensions

newupper a number, the rescaled upper bound for all dimensions

Value

z , inverse rescaling of vector x

See Also

[forwardRescale](#)

multiCOBRA *Perform multiple COBRA runs*

Description

Perform multiple COBRA runs. Each run starts with a different seed so that a different start point, a different initial design and different random restarts are choosen.

Usage

```
multiCOBRA(fn, lower, upper, nrun = 10, feval = 200, funcName = "GXX",
  fName = paste0("mult-", funcName, ".Rdata"), path = NULL, cobra = NULL,
  optim = NULL, target = 0.05, saveRdata = FALSE, ylim = c(1e-05,
  10000), plotPDF = FALSE, startSeed = 41)
```

Arguments

fn	objective function that is to be minimized, should return a vector of the objective function value and the constraint values
lower	lower bound of search space
upper	upper bound of search space
nrun	[10] number of runs
feval	[200] function evaluations per run
funcName	["GXX"] name of the problem
fName	the results (dfAll and others) are saved to <fname>.Rdata (only if saveRdata==TRUE)
path	[NULL] optional path
cobra	[NULL] list with COBRA settings. If NULL, initialize cobra with a suitable call to <code>cobraInit</code> .
optim	[NULL] the true optimum (or best known value) of the problem (only for diagnostics)
target	[0.05] a single run meets the target, if the final error is smaller than target
saveRdata	[FALSE] if TRUE, save results (dfAll,optim,target,fName,funcName) on <fName>.Rdata
ylim	the y limits
plotPDF	[FALSE] if TRUE, plot not only to current graphics device but to <fName>.pdf as well
startSeed	[41] after each run the seed is incremented by 1, starting with startSeed

Details

Side effect: An error plot showing each run and the mean and median of all runs (see `multiRunPlot`). The results (dfAll and others) are saved to <fName>.Rdata.

Value

mres, a list containing

cobra	the settings and results from last run
dfAll	a data frame with a result summary for all runs (see below)
z	a vector containing for each run the ever-best feasible objective value
z2	a data frame containing for each run the minimum error (if <code>optim</code> is available)

The data frame dfAll contains one row per iteration with columns (among others)

ffc fitness function calls (i.e. the iterations `cobra$iter`)

fitVal true fitness function value

fitSur surrogate fitness function value

feas is current iterate feasible on the true constraints?

feval number of evaluations of the internal optimizer on the surrogate functions (NA if it is a repairInfeasible-step)
XI the DRC element used in the current iteration
everBestFeas the ever-best feasible fitness function value
run the number of the current run
X1,X2,... the solution in (original) input space

Author(s)

Wolfgang Konen, Samineh Bagheri, Cologne University of Applied Sciences

See Also

[multiRunPlot](#), [cobraPhaseII](#)

Examples

```
## solve G11 problem nrun times and plot the results of all nrun runs
nrun=4
feval=25

## Defining the constrained problem (G11)
fn <- function(x) {
  y<-x[1]*x[1]+((x[2]-1)^2)
  y<-as.numeric(y)

  g1 <- as.numeric(+ (x[2] - x[1]^2))

  return(c(objective=y, g1=g1))
}
funcName="G11"
lower<-c(-1,-1)
upper<-c(+1,+1)

## Initializing and running cobra
cobra <- cobraInit(xStart=c(0,0), fn=fn, fName=funcName, lower=lower, upper=upper,
  feval=feval, initDesPoints=3*2, DOSAC=1, cobraSeed=1)

mres <- multiCOBRA(fn,lower,upper,nrun=nrun,feval=feval,optim=0.75
  ,cobra=cobra,funcName=funcName
  ,ylim=c(1e-12,1e-0),plotPDF=FALSE,startSeed=42)

## There are two true solutions at
## solu1 = c(-sqrt(0.5),0.5) and solu2 = c(+sqrt(0.5),0.5)
## where the true optimum is f(solu1) = f(solu2) = -0.75
## The solution from SACOBRA is close to one of the true solutions:
print(getXbest(mres$cobra))
print(getFbest(mres$cobra))
print(mres$z2)
```

multiRunPlot

Plot the results from multiple COBRA runs.

Description

Plot for each run one black curve 'error vs. iterations' and aggregate the mean curve (red) and the median curve (green) of all runs. 'error' is the distance between the ever-best feasible value and optim.

Usage

```
multiRunPlot(dfAll, optim = NULL, fName = "multiRun", main = "",
             xlim = NULL, ylim = c(1e-05, 10000), ylog = TRUE, xlog = FALSE,
             target = 0.05, plotPDF = FALSE, subPDF = NULL,
             legendWhere = "topright", absErr = FALSE)
```

Arguments

dfAll	the data frame of all runs, obtained with <code>multiCOBRA</code> or loaded from .Rdata file
optim	[NULL] the true optimum (or best known value) of the problem (only for diagnostics). If <code>optim=NULL</code> , we plot instead of errors the ever-best feasible values.
fName	["multiRun"] the name of the .Rdata file, printed as subtitle
main	[""] the name of the problem (e.g. "G01 problem"), printed as title
xlim	the x limits
ylim	the y limits
ylog	[TRUE] logarithmic y-axis
xlog	[FALSE] logarithmic x-axis
target	[0.05] a single run meets the target, if the final error is smaller than target
plotPDF	[FALSE] if TRUE, plot to 'fName'.pdf
subPDF	[NULL] optional subdirectory where .pdf should go
legendWhere	["topright"]
absErr	[FALSE] if TRUE, plot <code>abs(error)</code> instead of error.

Details

Print some diagnostic information: final median & mean error, percentage of runs which meet the target (only if `optim` is available).

Value

z3, a vector containing for each run the ever-best feasible objective value

Author(s)

Wolfgang Konen, Samineh Bagheri, Cologne University of Applied Sciences

See Also

[multiRunPlot_2](#), [multiCOBRA](#), [cobraPhaseII](#)

multiRunPlot_2

Plot the results from multiple COBRA runs.

Description

Plot for each run one black curve 'error vs. iterations' and aggregate the mean curve (red) and the median curve (green) of all runs. DIFFERENCE to [multiRunPlot](#): 'error' is the distance of the ever-best feasible point in input space to the true solution solu.

Usage

```
multiRunPlot_2(dfAll, solu, fName = "multiRun", main = "", xlim = NULL,
  ylim = c(1e-05, 10000), ylog = TRUE, xlog = FALSE, target = 0.05,
  plotPDF = FALSE, subPDF = NULL, legendWhere = "topright",
  absErr = FALSE)
```

Arguments

dfAll	the data frame of all runs, obtained with multiCOBRA or loaded from .Rdata file
solu	the true solution in input space of the problem (only for diagnostics).
fName	["multiRun"] the name of the .Rdata file, printed as subtitle
main	[""] the name of the problem (e.g. "G01 problem"), printed as title
xlim	the x limits
ylim	the y limits
ylog	[TRUE] logarithmic y-axis
xlog	[FALSE] logarithmic x-axis
target	[0.05] a single run meets the target, if the final error is smaller than target
plotPDF	[FALSE] if TRUE, plot to <fName>.pdf
subPDF	[NULL] optional subdirectory where .pdf should go
legendWhere	["topright"]
absErr	[FALSE] if TRUE, plot abs(error) instead of error.

Details

Print some diagnostic information: final median & mean error, percentage of runs which meet the target (only if opt im is available).

Value

z3, a vector containing for each run the ever-best feasible objective value

Author(s)

Wolfgang Konen, Samineh Bagheri, Cologne University of Applied Sciences

See Also

[multiRunPlot](#), [multiCOBRA](#), [cobraPhaseII](#)

plog

Monotonic transform

Description

The function is introduced in [Regis 2014] and extended here by a parameter p_{shift} . It is used to squash functions with a large range into a smaller range.

Let $y' = (y - p_{shift})$:

$$plog(y) = \ln(1 + y'), \quad \text{if } y' \geq 0$$

$$plog(y) = -\ln(1 - y'), \quad \text{if } y' < 0$$

Usage

```
plog(y, pShift = 0)
```

Arguments

y	function argument
pShift	shift

Value

$plog(y)$

See Also

[plogReverse](#)

plogReverse *Inverse of plog*

Description

Inverse of [plog](#)

Usage

```
plogReverse(y, pShift = 0)
```

Arguments

y	function argument
pShift	shift

Value

$plog^{-1}(y)$

See Also

[plog](#)

predict.RBFinter *Apply cubic or Gaussian or MQ RBF interpolation*

Description

Apply cubic or Gaussian or MQ RBF interpolation to a set of new data points for $d > 1$.

Usage

```
## S3 method for class 'RBFinter'
predict(rbf.model, newdata, ...)
```

Arguments

rbf.model	trained RBF model (or set of models), see trainCubicRBF or trainGaussRBF
newdata	matrix or data frame with d columns. Each row contains a data point x_i , $i = 1, \dots, n$
...	(not used)

Value

vector of model responses $s(x_i)$, one element for each data point x_i

- or -

if `rbf.model` is a set of m models, a $(n \times m)$ -matrix containing in each row the response $s_j(x_i)$ of all models $j = 1, \dots, m$ to x_i

Author(s)

Wolfgang Konen (<wolfgang.konen@th-koeln.de>)

See Also

[trainCubicRBF](#), [trainGaussRBF](#), [interpRBF](#)

repairChootinan	<i>Repair an infeasible solution with the method of Chootinan.</i>
-----------------	--

Description

Implements the method of [Choo2006] Chootinan & Chen "Constraint handling in genetic algorithms using a gradient-based repair method", Computers & Operations Research 33 (2006), p. 2263.

Usage

```
repairChootinan(x, gReal, rbf.model, cobra, checkIt = FALSE)
```

Arguments

<code>x</code>	an infeasible solution vector \vec{x} of dimension d
<code>gReal</code>	a vector $(g_1(\vec{x}), \dots, g_m(\vec{x}), h_1(\vec{x}), \dots, h_r(\vec{x}))$ holding the real constraint values at \vec{x}
<code>rbf.model</code>	the constraint surrogate models
<code>cobra</code>	parameter list, we need here lower lower bounds of search region upper upper bounds of search region ri a list with all parameters for repairChootinan trueFuncForSurrogate if TRUE (only for diagnostics), use the true constraint functions instead of the constraint surrogate models <code>rbf.model</code> fn true functions, only needed in case of <code>trueFuncForSurrogate==TRUE</code>
<code>checkIt</code>	[FALSE] if TRUE, perform a check whether the returned solution is really feasible. Needs access to the true constraint function <code>conFunc</code>

Value

`z`, a vector of dimension d with a repaired (hopefully feasible) solution

Author(s)

Wolfgang Konen, Cologne University of Applied Sciences

See Also

[repairInfeasRI2](#), [cobraPhaseII](#)

repairInfeasRI2

Repair an infeasible solution with the method RI2

Description

If the solution \vec{x} is infeasible, i.e. if there is any i or any j such that

$$g_i(\vec{x}) > 0 \text{ or } |h_j(\vec{x})| - \text{currentEps} > 0$$

:

1. Estimate the gradient of the constraint surrogate function(s) (go a tiny step in each dimension in the direction of constraint increase).
2. Take `cobrarimmax` random realizations in the 'feasible parallelepiped' and select among them the best feasible solution, based on the surrogates,
3. Check whether the new solution is for every dimension in the bounds [`cobra$lower`, `cobra$upper`] of the search region. If not, set the gradient to 0 in these dimensions and re-iterate from step 2.

There is no guarantee but a good chance, that the returned solution z will be feasible.

Usage

```
repairInfeasRI2(x, gReal, rbf.model, cobra, checkIt = FALSE)
```

Arguments

<code>x</code>	an infeasible solution vector \vec{x} of dimension <code>d</code>
<code>gReal</code>	a vector $(g_1(\vec{x}), \dots, g_m(\vec{x}), h_1(\vec{x}), \dots, h_r(\vec{x}))$ holding the real constraint values at \vec{x}
<code>rbf.model</code>	the constraint surrogate models
<code>cobra</code>	parameter list, we need here <code>lower</code> lower bounds of search region <code>upper</code> upper bounds of search region <code>ri</code> a list with all parameters for <code>repairInfeasRI2</code> , see defaultRI <code>trueFuncForSurrogate</code> if TRUE (only for diagnostics), use the true constraint functions instead of the constraint surrogate models <code>rbf.model</code> <code>fn</code> true functions, only needed in case of <code>trueFuncForSurrogate==TRUE</code>
<code>checkIt</code>	[FALSE] if TRUE, perform a check whether the returned solution is really feasible. Needs access to the true constraint functions.

Details

For further details see [Koch15a] Koch, P.; Bagheri, S.; Konen, W. et al. "A New Repair Method For Constrained Optimization". Proc. 17th Genetic and Evolutionary Computation Conference (GECCO), 2015.

Value

z, a vector of dimension d with a repaired (hopefully feasible) solution

Author(s)

Wolfgang Konen, Cologne University of Applied Sciences

See Also

[repairChootinan](#), [cobraPhaseII](#)

rescaleWrapper	<i>Return a rescaled function</i>
----------------	-----------------------------------

Description

Return a rescaled function

Usage

```
rescaleWrapper(fn, lower, upper, dimension, newlower, newupper)
```

Arguments

fn	function with argument x to be rescaled
lower	a vector with lower bounds in original input space
upper	a vector with lower bounds in original input space
dimension	length of vector lower and upper
newlower	a number, the rescaled lower bound for all dimensions
newupper	a number, the rescaled upper bound for all dimensions

Value

newfn, rescaled version of function fn

See Also

[forwardRescale](#), [inverseRescale](#)

setOpts	<i>Merge the options from a partial list and the default list</i>
---------	---

Description

Merge the options from a partial list and the default list

Usage

```
setOpts(opts, defaultOpt)
```

Arguments

opts	a partial list of options
defaultOpt	a list with default values for every element

Value

a list combined from opts and defaultOpt where every available element in opts overrides the default. For the rest of the elements the value from defaultOpt is taken.
A warning is issued for every element appearing in opts but not in defaultOpt

Author(s)

Samineh Bagheri, Wolfgang Konen, Cologne University of Applied Sciences

See Also

[defaultRI](#), [defaultSAC](#), [defaultTR](#), [defaultEquMu](#)

startCobra	<i>Start COBRA (constraint-based optimization) phase I and/or phase II</i>
------------	--

Description

Start COBRA (constraint-based optimization) phase I and/or phase II for object cobra

Usage

```
startCobra(cobra)
```

Arguments

cobra	initialized COBRA object, i.e. the return value from cobraInit
-------	--

Value

cobra, an object of class COBRA

See Also

[cobraInit](#), [cobraPhaseI](#), [cobraPhaseII](#)

Examples

```
## solve G01 problem

## defining the constraint problem: G01
fn<-function(x){
  obj<- sum(5*x[1:4])-(5*sum(x[1:4]*x[1:4]))-(sum(x[5:13]))
  g1<- (2*x[1]+2*x[2]+x[10]+x[11] - 10)
  g2<- (2*x[1]+2*x[3]+x[10]+x[12] - 10)
  g3<- (2*x[2]+2*x[3]+x[11]+x[12] - 10)

  g4<- -8*x[1]+x[10]
  g5<- -8*x[2]+x[11]
  g6<- -8*x[3]+x[12]

  g7<- -2*x[4]-x[5]+x[10]
  g8<- -2*x[6]-x[7]+x[11]
  g9<- -2*x[8]-x[9]+x[12]

  res<-c(obj, g1 ,g2 , g3
        , g4 , g5 , g6
        , g7 , g8 , g9)
  return(res)
}
fName="G01"
d=13
lower=rep(0,d)
upper=c(rep(1,9),rep(100,3),1)
set.seed(1)
xStart<-runif(d,min=lower,max=upper)

## Initializing cobra
cobra <- cobraInit(xStart=xStart, fn=fn, fName=fName, lower=lower, upper=upper,
                  feval=55, seqFeval=400, initDesPoints=3*d, DOSAC=1, cobraSeed=1)

cobra <- startCobra(cobra)
## The true solution is at solu = c(rep(1,9),rep(3,3),1)
## where the optimum is f(solu) = optim = -15
## The solutions from SACOBRA is close to this:
print(getXbest(cobra))
print(getFbest(cobra))

## Plot the resulting error (best-so-far feasible optimizer result - true optimum)
## on a logarithmic scale:
```

```
optim = -15
plot(cobra$df$Best-optim, log="y", type="l", ylab="error", xlab="iteration", main=fName)
```

trainCubicRBF

Fit cubic RBF interpolation to training data X for d>1.

Description

The model at a point $z = (z_1, \dots, z_d)$ is fitted using n sample points x_1, \dots, x_n

$$s(z) = \lambda_1 * \Phi(\|z - x_1\|) + \dots + \lambda_n * \Phi(\|z - x_n\|) + c_0 + c_1 * z_1 + \dots + c_d * z_d$$

where $\Phi(r) = r^3$ denotes the cubic radial basis function. The coefficients $\lambda_1, \dots, \lambda_n, c_0, c_1, \dots, c_d$ are determined by this training procedure.

This is for the default case `squares==FALSE`. In case `squares==TRUE` there are d additional pure square terms and the model is

$$s_{sq}(z) = s(z) + c_{d+1} * z_1^2 + \dots + c_{d+d} * z_d^2$$

In case `ptail==FALSE` the polynomial tail (all coefficients c_i) is omitted completely.

Usage

```
trainCubicRBF(xp, U, ptail = TRUE, squares = FALSE, rho = 0,
  DEBUG2 = FALSE, width = NA)
```

Arguments

xp	n points x_i of dimension d are arranged in $(n \times d)$ matrix xp
U	vector of length n , containing samples $f(x_i)$ of the scalar function f to be fitted - or - $(n \times m)$ matrix, where each column $1, \dots, m$ contains one vector of samples $f_j(x_i)$ for the m 'th model, $j=1, \dots, m$
ptail	[TRUE] flag, see description
squares	[FALSE] flag, see description
rho	[0.0] experimental: 0: interpolating, >0, approximating (spline-like) Gaussian RBFs
DEBUG2	[FALSE] if TRUE, save M and rhs on return value
width	[NA] non relevant for the parameter-free cubic RBF

Details

The linear equation system is solved via SVD inversion. Near-zero elements in the diagonal matrix D are set to zero in D^{-1} . This is numerically stable for rank-deficient systems.

Value

rbf.model, an object of class RBFinter, which is basically a list with elements:

coef	(n+d+1 x m) matrix holding in column m the coefficients for the m'th model: $\lambda_1, \dots, \lambda_n, c_0, c_1, \dots, c_d$. In case squares==TRUE it is an (n+2d+1 x m) matrix holding additionally the coefficients c_{d+1}, \dots, c_{d+d} .
xp	matrix xp
d	size of the polynomial tail. If length(d)==0 it means no polynomial tail will be used for the model. In case of ptail==T && squares==F d will be dimension+1 and in case of ptail==T && squares==T d will be 2*dimension+1
npts	number n of points x_i
ptail	TRUE or FALSE (see description)
squares	TRUE or FALSE (see description)
type	"CUBIC"
width	NA, irrelevant for the parameter-free cubic RBF

Author(s)

Wolfgang Konen (<wolfgang.konen@th-koeln.de>), Samineh Bagheri (<samineh.bagheri@th-koeln.de>)

See Also

[trainGaussRBF](#), [trainMQRBF](#) [predict.RBFinter](#), [interpRBF](#)

trainGaussRBF

Fit Gaussian RBF model to training data for $d > 1$.

Description

The model for a point $z = (z_1, \dots, z_d)$ is fitted using n sample points x_1, \dots, x_n

$$s(z) = \lambda_1 * \Phi(\|z - x_1\|) + \dots + \lambda_n * \Phi(\|z - x_n\|) + c_0 + c_1 * z_1 + \dots + c_d * z_d$$

where $\Phi(r) = \exp(-r^2/(2 * \sigma^2))$ denotes the Gaussian radial basis function with width σ . The coefficients $\lambda_1, \dots, \lambda_n, c_0, c_1, \dots, c_d$ are determined by this training procedure.

This is for the default case squares==FALSE. In case squares==TRUE there are d additional pure square terms and the model is

$$s_{sq}(z) = s(z) + c_{d+1} * z_1^2 + \dots + c_{d+d} * z_d^2$$

In case ptail==FALSE the polynomial tail (all coefficients c_i) is omitted completely.

The linear equation system is solved via SVD inversion. Near-zero elements in the diagonal matrix D are set to zero in D^{-1} . This makes rank-deficient systems numerically stable.

Usage

```
trainGaussRBF(xp, U, ptail = TRUE, squares = FALSE, width, RULE = "One",
              widthFactor = 1, rho = 0, DEBUG2 = F)
```

Arguments

xp	n points x_i of dimension d are arranged in (n x d) matrix xp
U	vector of length n, containing samples $u(x_i)$ of the scalar function u to be fitted - or - (n x m) matrix, where each column 1,...,m contains one vector of samples $u_j(x_i)$ for the m'th model, j=1,...,m
ptail	[TRUE] flag, see description
squares	[FALSE] flag, see 'Description'
width	[-1] either a positive real value which is the constant width σ for all Gaussians in all iterations, or -1. If -1, the appropriate width σ is calculated anew in each iteration with one of the rules RULE, based on the distribution of data points xp.
RULE	["One"] one out of ["One" "Two" "Three"], different rules for automatic estimation of width σ . Only relevant if width = -1,
widthFactor	[1.0] additional constant factor applied to each width σ
rho	[0.0] experimental: 0.0: interpolating, >0.0, approximating (spline-like) Gaussian RBFs
DEBUG2	[FALSE] if TRUE, save M and rhs on return value

Value

rbf.model, an object of class RBFinter, which is basically a list with elements:

coef	(n+d+1 x m) matrix holding in column m the coefficients for the m'th model: $\lambda_1, \dots, \lambda_n, c_0, c_1, \dots, c_d$. In case squares==TRUE it is an (n+2d+1 x m) matrix holding additionally the coefficients c_{d+1}, \dots, c_{d+d} .
xp	matrix xp
d	size of the polynomial tail. If length(d)==0 it means no polynomial tail will be used for the model. In case of ptail==T && squares==F d will be dimension+1 and in case of ptail==T && squares==T d will be 2*dimension+1
npts	number n of points x_i
ptail	TRUE or FALSE (see description)
squares	TRUE or FALSE (see description)
width	the calculated width σ
type	"GAUSS"

Author(s)

Wolfgang Konen, Samineh Bagheri

See Also

[trainCubicRBF](#), [predict.RBFinter](#), [interpRBF](#)

trainMQRBF

Fit multiquadric RBF model to training data for $d > 1$.

Description

The model for a point $z = (z_1, \dots, z_d)$ is fitted using n sample points x_1, \dots, x_n

$$s(z) = \lambda_1 * \Phi(\|z - x_1\|) + \dots + \lambda_n * \Phi(\|z - x_n\|) + c_0 + c_1 * z_1 + \dots + c_d * z_d$$

where $\Phi(r) = \sqrt{(1 + (r/\sigma)^2)}$ denotes the multiquadrics radial basis function with width σ . The coefficients $\lambda_1, \dots, \lambda_n, c_0, c_1, \dots, c_d$ are determined by this training procedure.

This is for the default case `squares==FALSE`. In case `squares==TRUE` there are d additional pure square terms and the model is

$$s_{sq}(z) = s(z) + c_{d+1} * z_1^2 + \dots + c_{d+d} * z_d^2$$

In case `ptail==FALSE` the polynomial tail (all coefficients c_i) is omitted completely.

Usage

```
trainMQRBF(xp, U, ptail = TRUE, squares = FALSE, width, RULE = "One",
widthFactor = 1, rho = 0, DEBUG2 = F)
```

Arguments

xp	n points x_i of dimension d are arranged in $(n \times d)$ matrix xp
U	vector of length n , containing samples $u(x_i)$ of the scalar function u to be fitted - or - $(n \times m)$ matrix, where each column $1, \dots, m$ contains one vector of samples $u_j(x_i)$ for the m 'th model, $j=1, \dots, m$
ptail	[TRUE] flag, see description
squares	[FALSE] flag, see 'Description'
width	[-1] either a positive real value which is the constant width σ for all Gaussians in all iterations, or -1. If -1, the appropriate width σ is calculated anew in each iteration with one of the rules RULE, based on the distribution of data points xp.
RULE	["One"] one out of ["One" "Two" "Three"], different rules for automatic estimation of width σ . Only relevant if <code>width = -1</code> ,
widthFactor	[1.0] additional constant factor applied to each width σ
rho	[0.0] experimental: 0.0: interpolating, >0.0, approximating (spline-like) Gaussian RBFs
DEBUG2	[FALSE] if TRUE, save M and rhs on return value

Details

The linear equation system is solved via SVD inversion. Near-zero elements in the diagonal matrix D are set to zero in D^{-1} . This makes rank-deficient systems numerically stable.

Value

`rbf.model`, an object of class `RBFinter`, which is basically a list with elements:

<code>coef</code>	($n+d+1 \times m$) matrix holding in column m the coefficients for the m 'th model: $\lambda_1, \dots, \lambda_n, c_0, c_1, \dots, c_d$. In case <code>squares==TRUE</code> it is an ($n+2d+1 \times m$) matrix holding additionally the coefficients c_{d+1}, \dots, c_{d+d} .
<code>xp</code>	matrix <code>xp</code>
<code>d</code>	size of the polynomial tail. If <code>length(d)==0</code> it means no polynomial tail will be used for the model. In case of <code>ptail==T && squares==F</code> <code>d</code> will be <code>dimension+1</code> and in case of <code>ptail==T && squares==T</code> <code>d</code> will be <code>2*dimension+1</code>
<code>npts</code>	number <code>n</code> of points x_i
<code>ptail</code>	TRUE or FALSE (see description)
<code>squares</code>	TRUE or FALSE (see description)
<code>width</code>	the calculated width σ
<code>type</code>	"MQ"

Author(s)

Wolfgang Konen, Samineh Bagheri

See Also

[trainCubicRBF](#), [predict.RBFinter](#), [interpRBF](#)

<code>trustRegion</code>	<i>Performs trust region refinement</i>
--------------------------	---

Description

If `cobra$TrustRegion==TRUE` (see [cobraInit](#)), then the `trustRegion` functionality is applied every iteration in order to refine the best solution so far. This function builds a local model around the best solution and runs a local search in the trust region to refine the best solution and find a better solution in the neighborhood.

Usage

```
trustRegion(cobra, center = cobra$xbest)
```

Arguments

cobra an object of class cobra, which is basically a list (see [cobraInit](#))
center [cobra\$xbest] the center of the trust region

Value

the modified cobra with new/updated elements

TRDONE logical, is TRUE if there are more than $d+1$ points in the trusted region and thus surrogates can be trained. Otherwise FALSE.

trustregX if TRDONE==TRUE the refined solution from the trust-region call, otherwise NA

If TRDONE==TRUE the relevant lists and counters (A, Fres, df, ...) of cobra will be updated in [cobraPhaseII](#) as well.

Author(s)

Samineh Bagheri (<samineh.bagheri@th-koeln.de>)

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