Package ‘ecr’

March 8, 2023

Title Evolutionary Computation in R

Description Framework for building evolutionary algorithms for both single- and multi-objective continuous or discrete optimization problems. A set of predefined evolutionary building blocks and operators is included. Moreover, the user can easily set up custom objective functions, operators, building blocks and representations sticking to few conventions. The package allows both a black-box approach for standard tasks (plug-and-play style) and a much more flexible white-box approach where the evolutionary cycle is written by hand.

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BugReports https://github.com/jakobbossek/ecr2/issues

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Description

Consider a data frame with results of multi-objective stochastic optimizers on a set of problems from different categories/groups (say indicated by column “group”). Occasionally, it is useful to unite the results of several groups into a meta-group. The function `addUnionGroup` aids in generation of such a meta-group while function `addAllGroup` is a wrapper around the former which generates a union of all groups.

Usage

```r
addUnionGroup(df, col, group, values)
```

```r
addAllGroup(df, col, group = "all")
```

Arguments

- `df` [data.frame]
  - Data frame.
- `col` [character(1)]
  - Column name of group-column.
- `group` [character(1)]
  - Name for new group.
- `values` [character(1)]
  - Subset of values within the value range of column `col`. 

Index

addUnionGroup  Grouping helpers
approximateNadirPoint

Value

data.frame Modified data frame.

Examples

df = data.frame(
  perf = runif(5),
  stringsAsFactors = FALSE)

df2 = addUnionGroup(df, col = "group", group = "A", values = c("A1", "A2"))
df3 = addAllGroup(df, col = "group", group = "ALL")

approximateNadirPoint Reference point approximations.

Description

Helper functions to compute nadir or ideal point from sets of points, e.g., multiple approximation sets.

Usage

approximateNadirPoint(..., sets = NULL)

approximateIdealPoint(..., sets = NULL)

Arguments

... [matrix]
  Arbirary number of matrizes.

sets [list]
  List of matrizes. This is an alternative way of passing the sets. Can be used exclusively or combined with .

Value

numeric Reference point.

See Also

Other EMOA performance assessment tools: approximateRefPoints(), approximateRefSets(),
computeDominanceRanking(), emoaIndEps(), makeEMOAIndicator(), niceCellFormatter(), normalize(),
plotDistribution(), plotFront(), plotScatter2d(), plotScatter3d(), toLatex()
approximateRefPoints  Helper function to estimate reference points.

Description

E.g., for calculation of dominated hypervolume.

Usage

approximateRefPoints(df, obj.cols = c("f1", "f2"), offset = 0, as.df = FALSE)

Arguments

df  [data.frame]
Data frame with the required structure, i.e. the data frame must contain a problem column "prob" as well as objective column(s).

obj.cols  [character(>= 2)]
Column names of the objective functions. Default is c("f1", "f2"), i.e., the bi-objective case is assumed.

offset  [numeric(1)]
Offset added to reference points. Default is 0.

as.df  [logical(1)]
Should a data.frame be returned? Default is FALSE. In this case a named list is returned.

Value

list | data.frame

See Also

Other EMOA performance assessment tools: approximateNadirPoint(), approximateRefSets(), computeDominanceRanking(), emoaIndEps(), makeEMOAIndicator(), niceCellFormater(), normalize(), plotDistribution(), plotFront(), plotScatter2d(), plotScatter3d(), toLatex()

approximateRefSets  Helper function to estimate reference set(s).

Description

The function takes an data frame with columns at least specified by obj.cols and “prob”. The reference set for each unique problem in column “prob” is then obtained by combining all approximation sets generated by all considered algorithms for the corresponding problem and filtering the non-dominated solutions.
Usage

approximateRefSets(df, obj.cols, as.df = FALSE)

Arguments

  df [data.frame]
  Data frame with the required structure.

  obj.cols [character(>= 2)]
  Column names of the objective functions.

  as.df [logical(1)]
  Should a data.frame be returned? Default is FALSE. In this case a named list is returned.

Value

list | data.frame
  Named list of matrices (names are the problems) or data frame with columns obj.cols and “prob”.

See Also

Other EMOA performance assessment tools: approximateNadirPoint(), approximateRefPoints(),
computeDominanceRanking(), emoaIndEps(), makeEMOAIndicator(), niceCellFormatter(), normalize(),
plotDistribution(), plotFront(), plotScatter2d(), plotScatter3d(), toLatex()

asemoa

Implementation of the NSGA-II EMOA algorithm by Deb.

Description

The AS-EMOA, short for aspiration set evolutionary multi-objective algorithm aims to incorporate expert knowledge into multi-objective optimization [1]. The algorithm expects an aspiration set, i.e., a set of reference points. It then creates an approximation of the pareto front close to the aspiration set utilizing the average Hausdorff distance.

Usage

asemoa(
  fitness.fun,
  n.objectives = NULL,
  minimize = NULL,
  n.dim = NULL,
  lower = NULL,
  upper = NULL,
  mu = 10L,
  aspiration.set = NULL,
  normalize.fun = NULL,
dist.fun = computeEuclideanDistance,
    p = 1,
    parent.selector = setup(selSimple),
    mutator = setup(mutPolynomial, eta = 25, p = 0.2, lower = lower, upper = upper),
    recombinator = setup(recSBX, eta = 15, p = 0.7, lower = lower, upper = upper),
    terminators = list(stopOnIters(100L))
)

Arguments

    fitness.fun [function]
    The fitness function.

    n.objectives [integer(1)]
    Number of objectives of obj.fun. Optional if obj.fun is a benchmark function from package smoof.

    minimize [logical(n.objectives)]
    Logical vector with ith entry TRUE if the ith objective of fitness.fun shall be minimized. If a single logical is passed, it is assumed to be valid for each objective.

    n.dim [integer(1)]
    Dimension of the decision space.

    lower [numeric]
    Vector of minimal values for each parameter of the decision space in case of float or permutation encoding. Optional if obj.fun is a benchmark function from package smoof.

    upper [numeric]
    Vector of maximal values for each parameter of the decision space in case of float or permutation encoding. Optional if obj.fun is a benchmark function from package smoof.

    mu [integer(1)]
    Population size. Default is 10.

    aspiration.set [matrix]
    The aspiration set. Each column contains one point of the set.

    normalize.fun [function]
    Function used to normalize fitness values of the individuals before computation of the average Hausdorff distance. The function must have the formal arguments "set" and "aspiration.set". Default is NULL, i.e., no normalization at all.

    dist.fun [function]
    Distance function used internally by Hausdorff metric to compute distance between two points. Expects a single vector of coordinate-wise differences between points. Default is computeEuclideanDistance.

    p [numeric(1)]
    Parameter p for the average Hausdorff metric. Default is 1.

    parent.selector [ecr_selector]
    Selection operator which implements a procedure to copy individuals from a given population to the mating pool, i.e., allow them to become parents.
mutator [ecr_mutator]
Mutation operator of type ecr_mutator.

recombinator [ecr_recombinator]
Recombination operator of type ecr_recombinator.

terminators [list]
List of stopping conditions of type “ecr_terminator”. Default is to stop after 100 iterations.

Value
ecr_multi_objective_result

Note
This is a pure R implementation of the AS-EMOA algorithm. It hides the regular ecr interface and offers a more R like interface while still being quite adaptable.

References

categorize

Description
Given a data frame and a grouping column of type factor or character this function generates a new grouping column which groups the groups.

Usage
categorize(df, col, categories, cat.col, keep = TRUE, overwrite = FALSE)

Arguments
df [data.frame]
Data frame.
col [character(1)]
Column name of group variable.
categories [list]
Named list. Names indicate the name of the category while the values are character vectors of values within the range of the col column.
computeAverageHausdorffDistance

**Value**

```r
```

**Description**

Computes the average Hausdroff distance measure between two point sets.

**Usage**

```r
computeAverageHausdorffDistance(
  A,
  B,
  p = 1,
  normalize = FALSE,
  dist.fun = computeEuclideanDistance
)
```

**Arguments**

- `A`  
  [matrix]
  First point set (each column corresponds to a point).

- `B`  
  [matrix]
  Second point set (each column corresponds to a point).

- `p`  
  [numeric(1)]
  Parameter p of the average Hausdorff metric. Default is 1.

- `normalize`  
  [logical(1)]
  Should the front be normalized on basis of B? Default is FALSE.

- `dist.fun`  
  [matrix]
  Distance function to compute distance between points x and y. Expects a single numeric vector d with the coordinate-wise differences di = (xi - yi). Default is computeEuclideanDist.
computeCrowdingDistance

Value
numeric(1) Average Hausdorff distance of sets A and B.

Description
The crowding distance is a measure of spread of solutions in the approximation of the Pareto front. It is used, e.g., in the NSGA-II algorithm as a second selection criterion.

Usage
computeCrowdingDistance(x)

Arguments
x [matrix]
Numeric matrix with each column representing a point.

Value
numeric Vector of crowding distance values.

References

computeDistanceFromPointToSetOfPoints

Description
Helper to compute distance between a single point and a point set.

Usage
computeDistanceFromPointToSetOfPoints(
a, 
B, 
dist.fun = computeEuclideanDistance
)

computeDistanceFromPointToSetOfPoints

Description
Computes distance between a single point and set of points.
**computeDominanceRanking**

**Arguments**

- `a` [numeric(1)]
  Point given as a numeric vector.

- `B` [matrix]
  Point set (each column corresponds to a point).

- `dist.fun` [matrix]
  Distance function to compute distance between points x and y. Expects a single numeric vector d with the coordinate-wise differences di = (xi - yi). Default is `computeEuclideanDist`.

**Value**

numeric(1)

---

**computeDominanceRanking**

*Ranking of approximation sets.*

**Description**

Ranking is performed by merging all approximation sets over all algorithms and runs per instance. Next, each approximation set C is assigned a rank which is 1 plus the number of approximation sets that are better than C. A set D is better than C, if for each point x ∈ C there exists a point in y ∈ D which weakly dominates x. Thus, each approximation set is reduced to a number – its rank. This rank distribution may act for first comparison of multi-objective stochastic optimizers. See [1] for more details. This function makes use of `parallelMap` to parallelize the computation of dominance ranks.

**Usage**

`computeDominanceRanking(df, obj.cols)`

**Arguments**

- `df` [data.frame]
  Data frame with columns at least “prob”, “algorithm”, “repl” and column names specified via parameter `obj.cols`.

- `obj.cols` [character(>= 2)]
  Column names in `df` which store the objective function values.

**Value**

data.frame Reduced `df` with columns “prob”, “algorithm”, “repl” and “rank”.

Note

Since pairwise non-domination checks are performed over all algorithms and algorithm runs this function may take some time if the number of problems, algorithms and/or replications is high.

References


See Also

Other EMOA performance assessment tools: approximateNadirPoint(), approximateRefPoints(), approximateRefSets(), emoaIndEps(), makeEMOAIndicator(), niceCellFormatter(), normalize(), plotDistribution(), plotFront(), plotScatter2d(), plotScatter3d(), toLatex()


computeGenerationalDistance

Computes Generational Distance.

Description

Helper to compute the Generational Distance (GD) between two sets of points.

Usage

computeGenerationalDistance(
  A,
  B,
  p = 1,
  normalize = FALSE,
  dist.fun = computeEuclideanDistance
)

Arguments

A [matrix]
First point set (each column corresponds to a point).

B [matrix]
Second point set (each column corresponds to a point).

p [numeric(1)]
Parameter p of the average Hausdoff metric. Default is 1.

normalize [logical(1)]
Should the front be normalized on basis of B? Default is FALSE.

dist.fun [matrix]
Distance function to compute distance between points x and y. Expects a single numeric vector d with the coordinate-wise differences d_i = (x_i - y_i). Default is computeEuclideanDist.
computeHV

Functions for the calculation of the dominated hypervolume (contribution).

**Description**

The function `computeHV` computes the dominated hypervolume of a set of points given a reference set whereby `computeHVContr` computes the hypervolume contribution of each point.

If no reference point is given the nadir point of the set \( x \) is determined and a positive offset with default 1 is added. This is to ensure that the reference point dominates all of the points in the reference set.

**Usage**

```r
computeHV(x, ref.point = NULL, ...)  
computeHVContr(x, ref.point = NULL, offset = 1)
```

**Arguments**

- `ref.point` [numeric\|NULL] Reference point. Set to the maximum in each dimension by default if not provided.
- `...` [any] Not used at the moment.
- `offset` [numeric(1)] Offset to be added to each component of the reference point only in the case where no reference is provided and one is calculated automatically.

**Value**

`numeric(1)` Dominated hypervolume in the case of `computeHV` and the dominated hypervolume contributions for each point in the case of `computeHVContr`.

**Note**

: Keep in mind that this function assumes all objectives to be minimized. In case at least one objective is to be maximized the matrix \( x \) needs to be transformed accordingly in advance.
**computeIndicators**  
*Computation of EMOA performance indicators.*

**Description**

Given a data.frame of Pareto-front approximations for different sets of problems, algorithms and replications, the function computes sets of unary and binary EMOA performance indicators. This function makes use of `parallelMap` to parallelize the computation of indicators.

**Usage**

```r
computeIndicators(
  df,
  obj.cols = c("f1", "f2"),
  unary.inds = NULL,
  binary.inds = NULL,
  normalize = FALSE,
  offset = 0,
  ref.points = NULL,
  ref.sets = NULL
)
```

**Arguments**

- `df`  
  [data.frame]  
  Data frame with columns `obj.cols`, “prob”, “algorithm” and “repl”.

- `obj.cols`  
  [character(>= 2)]  
  Column names of the objective functions. Default is `c("f1", "f2")`, i.e., the bi-objective case is assumed.

- `unary.inds`  
  [list]  
  Named list of unary indicators which shall be calculated. Each component must be another list with mandatory argument `fun` (the function which calculates the indicator) and optional argument `pars` (a named list of parameters for `fun`). Function `fun` must have the signature “function(points, arg1, ..., argk, ...)”. The arguments “points” and “...” are mandatory, the remaining are optional. The names of the components on the first level are used for the column names of the output data.frame. Default is `list(HV = list(fun = computeHV))`, i.e., the dominated Hypervolume indicator.

- `binary.inds`  
  [list]  
  Named list of binary indicators which shall be applied for each algorithm combination. Parameter `binary.inds` needs the same structure as `unary.inds`. However, the function signature of `fun` is slightly different: “function(points1, points2, arg1, ..., argk, ...)”. See function `emoaIndEps` for an example. Default is `list(EPS = list(fun = emoaIndEps))`.

- `normalize`  
  [logical]  
  If `TRUE`, normalize the indicators. Default is `FALSE`.

- `offset`  
  [numeric]  
  Offset for normalization. Default is `0`.

- `ref.points`  
  [numeric]  
  Reference points for normalization. Default is `NULL`.

- `ref.sets`  
  [numeric]  
  Reference sets for normalization. Default is `NULL`.
computeInvertedGenerationalDistance

Helper to compute the Inverted Generational Distance (IGD) between two sets of points.

Usage

computeInvertedGenerationalDistance(
  A,
  B,
  p = 1,
  normalize = FALSE,
  dist.fun = computeEuclideanDistance
)
dominated

Arguments

A [matrix]  
First point set (each column corresponds to a point).

B [matrix]  
Second point set (each column corresponds to a point).

p [numeric(1)]  
Parameter p of the average Hausdoff metric. Default is 1.

normalize [logical(1)]  
Should the front be normalized on basis of B? Default is FALSE.

dist.fun [matrix]  
Distance function to compute distance between points x and y. Expects a single numeric vector d with the coordinate-wise differences di = (xi - yi). Default is computeEuclideanDist.

Value

numeric(1)

dominated

Check for pareto dominance.

Description

These functions take a numeric matrix x where each column corresponds to a point and return a logical vector. The i-th position of the latter is TRUE if the i-th point is dominated by at least one other point for dominated and FALSE for nonDominated.

Usage
dominated(x)
nondominated(x)

Arguments

x [matrix]  
Numeric (d x n) matrix where d is the number of objectives and n is the number of points.

Value

logical
dominates

_Dominance relation check._

**Description**

Check if a vector dominates another (`dominates`) or is dominated by another (`isDominated`). There are corresponding infix operators `dominates` and `isDominatedBy`.

**Usage**

```
dominates(x, y)
```

```
isDominated(x, y)
```

```
x %dominates% y
```

```
x %isDominatedBy% y
```

**Arguments**

x [numeric]
First vector.

y [numeric]
Second vector.

**Value**

`logical(1)`

---

`doNondominatedSorting` _Fast non-dominated sorting algorithm._

**Description**

Fast non-dominated sorting algorithm proposed by Deb. Non-dominated sorting expects a set of points and returns a set of non-dominated fronts. In short words this is done as follows: the non-dominated points of the entire set are determined and assigned rank 1. Afterwards all points with the current rank are removed, the rank is increased by one and the procedure starts again. This is done until the set is empty, i.e., each point is assigned a rank.

**Usage**

```
doNondominatedSorting(x)
```
Arguments

x [matrix]
Numeric matrix of points. Each column contains one point.

Value

list List with the following components

  ranks Integer vector of ranks of length ncol(x). The higher the rank, the higher the domination
  front the corresponding point is located on.

  dom.counter Integer vector of length ncol(x). The i-th element is the domination number of the
  i-th point.

Note

This procedure is the key survival selection of the famous NSGA-II multi-objective evolutionary
algorithm (see nsgea).

References

[1] Deb, K., Pratap, A., and Agarwal, S. A Fast and Elitist Multiobjective Genetic Algorithm:

Description

The most flexible way to setup evolutionary algorithms with ecr is by explicitly writing the evolution-
ary loop utilizing various ecr utility functions. However, in everyday life R users frequently need
optimize a single-objective R function. The ecr function thus provides a more R like interface
for single objective optimization similar to the interface of the optim function.

Usage

ecr(
  fitness.fun,
  minimize = NULL,
  n.objectives = NULL,
  n.dim = NULL,
  lower = NULL,
  upper = NULL,
  n.bits,
  representation,
  mu,
  lambda,
  perm = NULL,
p.recomb = 0.7,
p.mut = 0.3,
survival.strategy = "plus",
n.elite = 0L,
log.stats = list(fitness = list("min", "mean", "max")),
log.pop = FALSE,
monitor = NULL,
initial.solutions = NULL,
parent.selector = NULL,
survival.selector = NULL,
mutator = NULL,
recombinator = NULL,
terminators = list(stopOnIters(100L)),
  ...
)

Arguments

fitness.fun [function]
The fitness function.

minimize [logical(n.objectives)]
Logical vector with ith entry TRUE if the ith objective of fitness.fun shall be minimized. If a single logical is passed, it is assumed to be valid for each objective.

n.objectives [integer(1)]
Number of objectives of obj.fun. Optional if obj.fun is a benchmark function from package smoof.

n.dim [integer(1)]
Dimension of the decision space.

lower [numeric]
Vector of minimal values for each parameter of the decision space in case of float or permutation encoding. Optional if obj.fun is a benchmark function from package smoof.

upper [numeric]
Vector of maximal values for each parameter of the decision space in case of float or permutation encoding. Optional if obj.fun is a benchmark function from package smoof.

n.bits [integer(1)]
Number of bits to use for binary representation.

representation [character(1)]
Genotype representation of the parameters. Available are “binary”, “float”, “permutation” and “custom”.

mu [integer(1)]
Number of individuals in the population.

lambda [integer(1)]
Number of individuals generated in each generation.
perm [integer(1) | vector]
Either a single integer number. In this case the set is assumed to be 1:perm. Alternatively, a set, i.e., a vector of elements can be passed which should form each individual.

p.recomb [numeric(1)]
Probability of two parents to perform crossover. Default is 0.7.

p.mut [numeric(1)]
The probability that the mutation operator will be applied to a child. Refers only to the application of the mutation operator, not to the probability of mutating individual genes of the respective child. Default is 0.1.

survival.strategy [character(1)]
Determines the survival strategy used by the EA. Possible are “plus” for a classical (mu + lambda) strategy and “comma” for (mu, lambda). Default is “plus”.

n.elite [integer(1)]
Number of fittest individuals of the current generation that shall be copied to the next generation without changing. Keep in mind, that the algorithm does not care about this option if the survival.strategy is set to 'plus'. Default is 0.

log.stats [list]
(Named) list of scalar functions to compute statistics on the fitness values in each generation. See initLogger for more information. Default is to log fitness minimum, mean and maximum values.

log.pop [logical(1)]
Shall the entire population be saved in each generation? Default is FALSE.

monitor [function]
Monitoring function. Default is NULL, i.e. no monitoring.

initial.solutions [list]
List of individuals which should be placed in the initial population. If the number of passed individuals is lower than mu, the population will be filled up by individuals generated by the corresponding generator. Default is NULL, i.e., the entire population is generated by the population generator.

parent.selector [ecr_selector]
Selection operator which implements a procedure to copy individuals from a given population to the mating pool, i.e., allow them to become parents.

survival.selector [ecr_selector]
Selection operator which implements a procedure to extract individuals from a given set, which should survive and set up the next generation.

mutator [ecr_mutator]
Mutation operator of type ecr_mutator.

recombinator [ecr_recombinator]
Recombination operator of type ecr_recombinator.
### terminators

[**list**]

List of stopping conditions of type “ecr_terminator”. Default is to stop after 100 iterations.

... [**any**]

Further arguments passed down to fitness.fun.

### Value

**ecr_result**

### Examples

```r
fn = function(x) {
  sum(x^2)
}
lower = c(-5, -5); upper = c(5, 5)
res = ecr(fn, n.dim = 2L, n.objectives = 1L, lower = lower, upper = lower,
         representation = "float", mu = 20L, lambda = 10L,
         mutator = setup(mutGauss, lower = lower, upper = upper))
```

### ecr_parallelization

**Parallelization in ecr**

### Description

In ecr it is possible to parallelize the fitness function evaluation to make use, e.g., of multiple CP cores or nodes in a HPC cluster. For maximal flexibility this is realized by means of the **parallelMap** package (see the official GitHub page for instructions on how to set up parallelization). The different levels of parallelization can be specified in the parallelStart* function. At the moment only the level “ecr.evaluateFitness” is supported.

Keep in mind that parallelization comes along with some overhead. Thus activating parallelization, e.g., for evaluation a fitness function which is evaluated lightning-fast, may result in higher computation time. However, if the function evaluations are computationally more expensive, parallelization leads to significant running time benefits.

### ecr_result

**Result object.**

### Description

S3 object returned by **ecr** containing the best found parameter setting and value in the single-objective case and the Pareto-front/-set in case of a multi-objective optimization problem. Moreover a set of further information, e.g., reason of termination, the control object etc. are returned.

The single objective result object contains the following fields:
emoaIndEps

**task** The ecr_optimization_task.
**best.x** Overall best parameter setting.
**best.y** Overall best objective value.
**log** Logger object.
**last.population** Last population.
**last.fitness** Numeric vector of fitness values of the last population.
**message** Character string describing the reason of termination.

In case of a solved multi-objective function the result object contains the following fields:

**task** The ecr_optimization_task.
**log** Logger object.
**pareto.idx** Indizes of the non-dominated solutions in the last population.
**pareto.front** (n x d) matrix of the approximated non-dominated front where n is the number of non-dominated points and d is the number of objectives.
**pareto.set** Matrix of decision space values resulting with objective values given in pareto.front.
**last.population** Last population.
**message** Character string describing the reason of termination.

---

emoaIndEps  \hspace{1em} EMOA performance indicators

**Description**

Functions for the computation of unary and binary measures which are useful for the evaluation of the performance of EMOAs. See the references section for literature on these indicators.

Given a set of points points, emoaIndEps computes the unary epsilon-indicator provided a set of reference points ref.points.

The emoaIndHV function computes the hypervolume indicator Hyp(X, R, r). Given a set of points X (points), another set of reference points R (ref.points) (which maybe the true Pareto front) and a reference point r (ref.point) it is defined as Hyp(X, R, r) = HV(R, r) - HV(X, r).

Function emoaIndR1, emoaIndR2 and emoaIndR3 calculate the R1, R2 and R3 indicator respectively.

Function emoaIndMD computes the minimum distance indicator, i.e., the minimum Euclidean distance between two points of the set points while function emoaIndM1 determines the mean Euclidean distance between points and points from a reference set ref.points.

Function emoaIndC calculates the coverage of the sets points (A) and ref.points (B). This is the ratio of points in B which are dominated by at least one solution in A.

emoaIndONVG calculates the “Overall Non-dominated Vector Generation” indicator. Despite its complicated name it is just the number of non-dominated points in points.

Functions emoaIndSP and emoaIndDelta calculate spacing indicators. The former was proposed by Schott: first calculate the sum of squared distances between minimal distances of points to all other points and the mean of these minimal distance. Next, normalize by the number of points minus 1 and finally calculate the square root. In contrast, Delta-indicator
Usage

emoaIndEps(points, ref.points, ...)

emoaIndHV(points, ref.points, ref.point = NULL, ...)

emoaIndR1(
  points,
  ref.points,
  ideal.point = NULL,
  nadir.point = NULL,
  lambda = NULL,
  utility = "tschebycheff",
  ...
)

emoaIndR2(
  points,
  ref.points,
  ideal.point = NULL,
  nadir.point = NULL,
  lambda = NULL,
  utility = "tschebycheff",
  ...
)

emoaIndR3(
  points,
  ref.points,
  ideal.point = NULL,
  nadir.point = NULL,
  lambda = NULL,
  utility = "tschebycheff",
  ...
)

emoaIndMD(points, ...)

emoaIndC(points, ref.points, ...)

emoaIndM1(points, ref.points, ...)

emoaIndONVG(points, ...)

emoaIndGD(
  points,
  ref.points,
  p = 1,
  normalize = FALSE,
emoaIndEps

\[
\text{dist.fun} = \text{computeEuclideanDistance},
\]

)

emoaIndIGD(
  points,
  ref.points,
  p = 1,
  normalize = FALSE,
  dist.fun = \text{computeEuclideanDistance},
  ...
)

emoaIndDeltap(
  points,
  ref.points,
  p = 1,
  normalize = FALSE,
  dist.fun = \text{computeEuclideanDistance},
  ...
)

emoaIndSP(points, ...)

emoaIndDelta(points, ...)

Arguments

points [matrix]
Matrix of points.

ref.points [matrix]
Set of reference points.

... [any]
Not used at the moment.

ref.point [numeric]
A single reference point used, e.g., for the computation of the hypervolume indicator via emoaIndHV. If NULL the nadir point of the union of the points and ref.points is used.

ideal.point [numeric]
The utopia point of the true Pareto front, i.e., each component of the point contains the best value if the other objectives are neglected.

nadir.point [numeric]
Nadir point of the true Pareto front.

lambda [integer(1)]
Number of weight vectors to use in estimating the utility function.

utility [character(1)]
evaluateFitness

Name of the utility function to use. Must be one of “weightedsum”, “tschebycheff” or “augmented tschbycheff”.

p

[numeric(1)]

Parameter p of the average Hausdoff metric. Default is 1.

normalize

[logical(1)]

Should the front be normalized on basis of B? Default is FALSE.

dist.fun

[matrix]

Distance function to compute distance between points x and y. Expects a single numeric vector d with the coordinate-wise differences di = (xi - yi). Default is computeEuclideanDist.

Value

numeric(1) Epsilon indicator.

See Also

Other EMOA performance assessment tools: approximateNadirPoint(), approximateRefPoints(), approximateRefSets(), computeDominanceRanking(), makeEMOAIndicator(), niceCellFormater(), normalize(), plotDistribution(), plotFront(), plotScatter2d(), plotScatter3d(), toLatex()

evaluateFitness

Computes the fitness value(s) for each individual of a given set.

Description

This function expects a list of individuals, computes the fitness and always returns a matrix of fitness values; even in single-objective optimization a (1 x n) matrix is returned for consistency, where n is the number of individuals. This function makes use of parallelMap to parallelize the fitness evaluation.

Usage

evaluateFitness(control, inds, ...)

Arguments

control

[ecr_control]

Control object.

inds

[list]

List of individuals.

...

[any]

Optional parameters passed down to fitness function.

Value

matrix .
**Description**

Given a data frame and a column name, function `explode` splits the content of a column by a specified delimiter (thus exploded) into multiple columns. Function `implode` does vice versa, i.e., given a non-empty set of column names or numbers, the function glues together the columns. Hence, functions `explode` and `implode` are kind of inverse to each other.

**Usage**

```
explode(df, col, by = "." , keep = FALSE, col.names = NULL)
```

```
implode(df, cols, by = "." , keep = FALSE, col.name)
```

**Arguments**

- `df` [data.frame] Data frame.
- `col` [character(1)] Name of column which should be exploded.
- `by` [character(1)] Delimiter used to split cell entries (for `explode`) or glue them together (for `implode`).
- `keep` [logical(1)] Should exploded or imploded source column be kept? Default is `FALSE`.
- `col.names` [character] Names of new columns. Default is "col.1", ..., "col.k", where k is the number of elements each cell in column `col` is split into.
- `cols` [character(1)] Names of columns (or column number) which should be imploded.
- `col.name` [character(1)] Name of new column.

**Value**

data.frame Modified data frame.

**Examples**

```
df = data.frame(x = 1:3, y = c("a.c", "a.b", "a.c"))
df.ex = explode(df, col = "y", col.names = c("y1", "y2"))
df.im = implode(df.ex, cols = c("y1", "y2"), by = "---", col.name = "y", keep = TRUE)
```
filterDuplicated  
*Filter approximation sets by duplicate objective vectors.*

**Description**

Filter approximation sets by duplicate objective vectors.

**Usage**

```r
filterDuplicated(x, ...)  
```

### S3 method for class 'data.frame'

```r
filterDuplicated(x, ...)  
```

### S3 method for class 'matrix'

```r
filterDuplicated(x, ...)  
```

### S3 method for class 'ecr_multi_objective_result'

```r
filterDuplicated(x, ...)  
```

### S3 method for class 'list'

```r
filterDuplicated(x, ...)  
```

**Arguments**

- **x**  
  [object]
  Object of type data frame (objectives column-wise), matrix (objectives row-wise), `ecr_multi_objective_result` or list (with components “pareto.front”) and “pareto.set”.

- **...**  
  [any]
  Not used at the moment

**Value**

- **object**  
  Modified input `x`.

**Note**

Note that this may be misleading if there can be solutions with identical objective function values but different values in decision space.
generateOffspring  

**Helper functions for offspring generation**

**Description**

Function `mutate` expects a control object, a list of individuals, and a mutation probability. The mutation operator registered in the control object is then applied with the given probability to each individual. Function `recombinate` expects a control object, a list of individuals as well as their fitness matrix and creates `lambda` offspring individuals by recombining parents from `inds`. Which parents take place in the parent selection depends on the `parent.selector` registered in the control object. Finally, function `generateOffspring` is a wrapper for both `recombinate` and `mutate`. Both functions are applied subsequently to generate new individuals by variation and mutation.

**Usage**

```r
generateOffspring(control, inds, fitness, lambda, p.recomb = 0.7, p.mut = 0.1)
mutate(control, inds, p.mut = 0.1, slot = "mutate", ...)
recombinate(
  control,
  inds,
  fitness,
  lambda = length(inds),
  p.recomb = 0.7,
  slot = "recombine",
  ...
)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>control</code></td>
<td>[ecr_control] Control object.</td>
</tr>
<tr>
<td><code>inds</code></td>
<td>[list] List of individuals.</td>
</tr>
<tr>
<td><code>fitness</code></td>
<td>[matrix] Matrix of fitness values (each column contains the fitness value(s) of one individual).</td>
</tr>
<tr>
<td><code>lambda</code></td>
<td>[integer(1)] Number of individuals generated in each generation.</td>
</tr>
<tr>
<td><code>p.recomb</code></td>
<td>[numeric(1)] Probability of two parents to perform crossover. Default is 0.7.</td>
</tr>
<tr>
<td><code>p.mut</code></td>
<td>[numeric(1)] The probability that the mutation operator will be applied to a child. Refers only to the application of the mutation operator, not to the probability of mutating individual genes of the respective child. Default is 0.1.</td>
</tr>
</tbody>
</table>
The slot of the control object which contains the registered operator to use. Default is “mutate” for mutate and “recombine” for recombinator. In most cases there is no need to change this. However, it might be useful if you make use of different mutation operators registered, e.g., in the slots “mutate1” and “mutate2”.

Further arguments passed down to recombinator/mutator. There parameters will overwrite parameters in par.list.

Value

list  List of individuals.

generatesMultipleChildren

Description

$Does the recombinator generate multiple children?$

Usage

generatesMultipleChildren(recombinator)

Arguments

recombinator  [function]

Actual mutation operator.

Value

logical  Boolean

generators

Description

Utility functions to build a set of individuals. The function gen expects an R expression and a number n in order to create a list of n individuals based on the given expression. Functions genBin, genPerm and genReal are shortcuts for initializing populations of binary strings, permutations or real-valued vectors respectively.
**Usage**

```r
gen(expr, n)
genBin(n, n.dim)
genPerm(n, n.dim)
genReal(n, n.dim, lower, upper)
```

**Arguments**

- `expr` [R expression]
  Expression to generate a single individual.
- `n` [integer(1)]
  Number of individuals to create.
- `n.dim` [integer(1)]
  Dimension of the decision space.
- `lower` [numeric]
  Vector of minimal values for each parameter of the decision space in case of float encoding.
- `upper` [numeric]
  Vector of maximal values for each parameter of the decision space in case of float encoding.

**Value**

`list`

---

**getFront**

*Extract fitness values from Pareto archive.*

**Description**

Get all non-dominated points in objective space, i.e., an \((m \times n)\) matrix of fitness with \(m\) being the number of objectives and \(n\) being the number of non-dominated points in the Pareto archive.

**Usage**

```r
getFront(x)
```

**Arguments**

- `x` [ecr_pareto_archive]
  Pareto archive.

**Value**

`matrix`
getIndividuals

Extract individuals from Pareto archive.

Description
Get the non-dominated individuals logged in the Pareto archive.

Usage
getIndividuals(x)

Arguments
x [ecr_pareto_archive]
Pareto archive.

Value
list

See Also
Other ParetoArchive: getSize(), initParetoArchive(), updateParetoArchive()

getNumberOfChildren

Number of children

Description
Returns the number children generated by the recombinator

Usage
getNumberOfChildren(recombinator)

Arguments
recombinator [function]
Actual mutation operator.

Value
numeric Number of children generated
getNumberOfParentsNeededForMating

Number of parents needed for mating

Description

Returns the number of parents needed for mating.

Usage

getNumberOfParentsNeededForMating(recombinator)

Arguments

recombinator [function]
Actual mutation operator.

Value

numeric Number of Parents need for mating

getPopulationFitness

Access to logged population fitness.

Description

Returns the fitness values of all individuals as a data.frame with columns f1, ..., fo, where o is the number of objectives and column “gen” for generation.

Usage

getPopulationFitness(log, trim = TRUE)

Arguments

log [ecr_logger]
The log generated by initLogger.
trim [logical(1)]
Should unused lines in the logged be cut off? Usually one wants this behaviour. Thus the default is TRUE.

Value

list List of populations.

See Also

Other logging: getPopulations(), getStatistics(), initLogger(), updateLogger()
getPopulations

Access to logged populations.

Description

Simple getter for the logged populations.

Usage

getPopulations(log, trim = TRUE)

Arguments

log [ecr_logger]
The log generated by initLogger.

trim [logical(1)]
Should unused lines in the logged be cut off? Usually one wants this behaviour. Thus the default is TRUE.

Details

This function throws an error if the logger was initialized with log.pop = FALSE (see initLogger).

Value

list List of populations.

See Also

Other logging: getPopulationFitness(), getStatistics(), initLogger(), updateLogger()
**getStatistics**

**Value**

integer(1)

**See Also**

Other ParetoArchive: `getIndividuals()`, `initParetoArchive()`, `updateParetoArchive()`

---

**getStatistics**

*Access the logged statistics.*

**Description**

Simple getter for the logged fitness statistics.

**Usage**

```r
getStatistics(log, trim = TRUE)
```

**Arguments**

- `log` [ecr_logger]
  - The log generated by `initLogger`.
- `trim` [logical(1)]
  - Should unused lines in the logged be cut off? Usually one wants this behaviour. Thus the default is TRUE.

**Value**

data.frame  Logged statistics.

**See Also**

Other logging: `getPopulationFitness()`, `getPopulations()`, `initLogger()`, `updateLogger()`
**getSupportedRepresentations**

*Get supported representations.*

**Description**

Returns the character vector of representation which the operator supports.

**Usage**

```r
getSupportedRepresentations(operator)
```

**Arguments**

- `operator`  
  `ecr_operator`  
  Operator object.

**Value**

- character  
  Vector of representation types.

---

**initECRControl**

*Control object generator.*

**Description**

The control object keeps information on the objective function and a set of evolutionary components, i.e., operators.

**Usage**

```r
initECRControl(fitness.fun, n.objectives = NULL, minimize = NULL)
```

**Arguments**

- `fitness.fun`  
  `function`  
  The fitness function.

- `n.objectives`  
  `integer(1)`  
  Number of objectives of `obj.fun`. Optional if `obj.fun` is a benchmark function from package `smoof`.

- `minimize`  
  `logical(n.objectives)`  
  Logical vector with ith entry TRUE if the ith objective of `fitness.fun` shall be minimized. If a single logical is passed, it is assumed to be valid for each objective.
**Description**

Logging is a central aspect of each EA. Besides the final solution(s) especially in research often we need to keep track of different aspects of the evolutionary process, e.g., fitness statistics. The logger of ecr keeps track of different user-defined statistics and the population. It may also be used to check stopping conditions (see `makeECRTerminator`). Most important this logger is used internally by the `ecr` black-box interface.

**Usage**

```r
initLogger(
  control, 
  log.stats = list(fitness = list("min", "mean", "max")),
  log.extras = NULL,
  log.pop = FALSE,
  init.size = 1000L
)
```

**Arguments**

- `control` [ecr_control]
  Control object.

- `log.stats` [list]
  List of lists for statistic computation on attributes of the individuals of the population. Each entry should be named by the attribute it should be based on, e.g., fitness, and should contain a list of R functions as a character string or a list with elements `fun` for the function, and `pars` for additional parameters which shall be passed to the corresponding function. Each function is required to return a scalar numeric value. By default the minimum, mean and maximum of the fitness values is computed. Since fitness statistics are the most important ones these do not have to be stored as attributes, but can be passed as a matrix to the update function.

- `log.extras` [character]
  Possibility to instruct the logger to store additional scalar values in each generation. Named character vector where the names indicate the value to store and the value indicates the corresponding data types. Currently we support all atomic modes of `vector` except “factor” and “raw”.

- `log.pop` [logical(1)]
  Shall the entire population be saved in each generation? Default is `FALSE`.  

**Value**

`ecr_control`
init.size [integer(1)]

Initial number of rows of the slot of the logger, where the fitness statistics are stored. The size of the statistics log is doubled each time an overflow occurs. Default is 1000.

Value
ecr_logger An S3 object of class ecr_logger with the following components:

- log.stats The log.stats list.
- log.pop The log.pop parameter.
- init.size Initial size of the log.
- env The actual log. This is an R environment which ensures, that in-place modification is possible.

Note
Statistics are logged in a data.frame.

See Also
Other logging: getPopulationFitness(), getPopulations(), getStatistics(), updateLogger()

Examples

```r
control = initECRControl(function(x) sum(x), minimize = TRUE,
                           n.objectives = 1L)
control = registerECROperator(control, "mutate", mutBitflip, p = 0.1)
control = registerECROperator(control, "selectForMating", selTournament, k = 2)
control = registerECROperator(control, "selectForSurvival", selGreedy)

log = initLogger(control,
                  log.stats = list(
                      fitness = list("mean", "myRange" = function(x) max(x) - min(x)),
                      age = list("min", "max")
                  ),
                  log.pop = TRUE, init.size = 1000L)

# simply pass stuff down to control object constructor
population = initPopulation(mu = 10L, genBin, n.dim = 10L)
fitness = evaluateFitness(control, population)

# append fitness to individuals and init age
for (i in seq_along(population)) {
  attr(population[[i]], "fitness") = fitness[, i]
  attr(population[[i]], "age") = 1L
}

for (iter in seq_len(10)) {
  # generate offspring
  offspring = generateOffspring(control, population, fitness, lambda = 5)
  fitness.offspring = evaluateFitness(control, offspring)
```
# update age of population
for (i in seq_along(population)) {
  attr(population[[i]], "age") = attr(population[[i]], "age") + 1L
}

# set offspring attributes
for (i in seq_along(offspring)) {
  attr(offspring[[i]], "fitness") = fitness.offspring[, i]
  # update age
  attr(offspring[[i]], "age") = 1L
}

sel = replaceMuPlusLambda(control, population, offspring)

population = sel$population
fitness = sel$fitness

# do some logging
updateLogger(log, population, n.evals = 5)
}
head(getStatistics(log))

initParetoArchive

Initialize Pareto Archive.

Description

A Pareto archive is usually used to store all / a part of the non-dominated points stored during a run of an multi-objective evolutionary algorithm.

Usage

initParetoArchive(control, max.size = Inf, trunc.fun = NULL)

Arguments

control [ecr_control]
  Control object.

max.size [integer(1)]
  Maximum capacity of the Pareto archive, i.e., the maximal number of non-dominated points which can be stored in the archive. Default is Inf, i.e., (theoretically) unbounded capacity.

trunc.fun [function(archive, inds, fitness, ...)]
  In case the archive is limited in capacity, i.e., max.size is not infinite, this function is called internally if an archive overflow occurs. This function expects the archive, a list of individuals inds, a matrix of fitness values (each column contains the fitness value(s) of one individual) fitness and further optional arguments ... which may be used by the internals of trunc.fun. The function must return a list with components “fitness” and “inds” which shall be the subsets of fitness and inds respectively, which should be kept by the archive.
initPopulation

Value
ecr_pareto_archive

See Also
Other ParetoArchive: getIndividuals(), getSize(), updateParetoArchive()

initPopulation

Helper function to build initial population.

Description
Generates the initial population. Optionally a set of initial solutions can be passed.

Usage
initPopulation(mu, gen.fun, initial.solutions = NULL, ...)

Arguments
mu [integer(1)]
Number of individuals in the population.

gen.fun [function]
Function used to generate initial solutions, e.g., genBin.

initial.solutions [list]
List of individuals which should be placed in the initial population. If the number of passed individuals is lower than mu, the population will be filled up by individuals generated by the corresponding generator. Default is NULL, i.e., the entire population is generated by the population generator.

... [any]
Further parameters passed to gen.fun.

Value
ecr_population
is.supported

Description

Check if the given operator supports a certain representation, e.g., “float”.

Usage

is.supported(operator, representation)

Arguments

operator [ecr_operator]
Object of type ecr_operator.

representation [character(1)]
Representation as a string.

Value

logical(1) TRUE, if operator supports the representation type.

isEcrOperator

Description

Checks if the passed object is of type ecr_operator.

Usage

isEcrOperator(obj)

Arguments

obj [any]
Arbitrary R object to check.

Value

logical(1)
makeECRMonitor  
Factory method for monitor objects.

Description
Monitor objects serve for monitoring the optimization process, e.g., printing some status messages to the console. Each monitor includes the functions before, step and after, each being a function and expecting a log log of type ecr_logger and ... as the only parameters. This way the logger has access to the entire log.

Usage
makeECRMonitor(before = NULL, step = NULL, after = NULL, ...)

Arguments
before [function]
Function called one time after initialization of the EA.

step [function]
Function applied after each iteration of the algorithm.

after [function]
Function applied after the EA terminated.

... [any]
Not used.

Value
ecr_monitor  Monitor object.

makeEMOAIndicator  
Constructor for EMOA indicators.

Description
Simple wrapper for function which compute performance indicators for multi-objective stochastic algorithm. Basically this function appends some meta information to the passed function fun.

Usage
makeEMOAIndicator(fun, minimize, name, latex.name)
makeMutator

Arguments

fun [function(points, ...)]
Function which expects a numeric matrix “points” as first argument. Optional named arguments (often “ref.point” for a reference point or “ref.points” for a reference set, e.g., the true Pareto-front) are allowed. See implementations of existing indicators for examples.

minimize [logical(1)]
Lower values indicate better performance?

name [character(1)]
Short name of the indicator. Used, e.g., as column name for the indicator in the data.frame returned by `computeIndicators`.

latex.name [character(1)]
LaTeX representation of the indicator. Used in LaTeX-table output statistical tests (see `toLatex`).

Value

function(points, ...) Argument fun with all other arguments appended.

See Also

Other EMOA performance assessment tools: `approximateNadirPoint()`, `approximateRefPoints()`, `approximateRefSets()`, `computeDominanceRanking()`, `emoaIndEps()`, `niceCellFormatter()`, `normalize()`, `plotDistribution()`, `plotFront()`, `plotScatter2d()`, `plotScatter3d()`, `toLatex()`

---

Description

Helper function which constructs a mutator, i.e., a mutation operator.

Usage

`makeMutator(mutator, supported = getAvailableRepresentations())`

Arguments

mutator [function]
Actual mutation operator.

supported [character]
Vector of strings/names of supported parameter representations. Possible choices: “permutation”, “float”, “binary” or “custom”.

Value

ecr_mutator Mutator object.
**makeOperator**  
_Construct evolutionary operator._

**Description**  
Helper function which constructs an evolutionary operator.

**Usage**

```
makeOperator(operator, supported = getAvailableRepresentations())
```

**Arguments**

- **operator**  
  [function]  
  Actual operator.

- **supported**  
  [character]  
  Vector of names of supported parameter representations. Possible choices: “per-mutation”, “float”, “binary” or “custom”.

**Value**

ecr_operator  
Operator object.

**Note**

In general you will not need this function, but rather one of its derivatives like _makeMutator_ or _makeSelector_.

---

**makeOptimizationTask**  
_Creates an optimization task._

**Description**

An optimization task consists of the fitness/objective function, the number of objectives, the “direction” of optimization, i.e., which objectives should be minimized/maximized and the names of the objectives.

**Usage**

```
makeOptimizationTask(
  fun,
  n.objectives = NULL,
  minimize = NULL,
  objective.names = NULL
)
```
**Arguments**

- **fun** [function | smoof_function]
  Fitness/objective function.

- **n.objectives** [integer(1)]
  Number of objectives. This must be a positive integer value unless `fun` is of type smoof_function.

- **minimize** [logical]
  A logical vector indicating which objectives to minimize/maximize. By default all objectives are assumed to be minimized.

- **objective.names** [character]
  Names for the objectives. Default is NULL. In this case the names are set to y1, ..., yn with n equal to n.objectives and simply y in the single-objective case.

**Value**

ecr_optimization_task

---

**makeRecombinator**  
*Construct a recombination operator.*

**Description**

Helper function which constructs a recombinator, i.e., a recombination operator.

**Usage**

```r
makeRecombinator(
  recombinator,
  supported = getAvailableRepresentations(),
  n.parents = 2L,
  n.children = NULL
)
```

**Arguments**

- **recombinator** [function]
  Actual mutation operator.

- **supported** [character]
  Vector of strings/names of supported parameter representations. Possible choices: "permutation", "float", "binary" or "custom".

- **n.parents** [integer(1)]
  Number of parents supported.

- **n.children** [integer(1)]
  How many children does the recombinator produce? Default is 1.
**makeSelector**

Value

ecr_recombinator  Recombinator object.

Note

If a recombinator returns more than one child, the `multiple.children` parameter needs to be `TRUE`, which is the default. In case of multiple children produced these have to be placed within a list.

---

**Description**

Helper function which defines a selector method, i.e., an operator which takes the population and returns a part of it for mating or survival.

**Usage**

```r
makeSelector(
  selector,
  supported = getAvailableRepresentations(),
  supported.objectives,
  supported.opt.direction = "minimize"
)
```

**Arguments**

- **selector**  [function]  Actual selection operator.
- **supported**  [character]  Vector of strings/names of supported parameter representations. Possible choices: “permutation”, “float”, “binary” or “custom”.
- **supported.objectives**  [character]  At least one of “single-objective” or “multi-objective”.
- **supported.opt.direction**  [character(1-2)]  Does the selector work for maximization tasks xor minimization tasks or both? Default is “minimize”, which means that the selector selects in favour of low fitness values.

**Value**

ecr_selector  Selector object.
makeTerminator

Generate stopping condition.

Description

Wrap a function within a stopping condition object.

Usage

makeTerminator(condition.fun, name, message)

Arguments

condition.fun [function]
Function which takes a logger object log (see initLogger) and returns a single logical.

name [character(1)]
Identifier for the stopping condition.

message [character(1)]
Message which should be stored in the termination object, if the stopping condition is met.

Value

ecr_terminator

mcMST

mcMST

Description

Pareto-front approximations for some graph problems obtained by several algorithms for the multi-criteria minimum spanning tree (mcMST) problem.

Usage

mcMST
Format

A data frame with four variables:

f1 First objective (to be minimized).
f2 Second objective (to be minimized).
algorithm Short name of algorithm used.
prob Short name of problem instance.
repl Algorithm run.

The data is based on the mcMST package.

---

mutBitflip Bitflip mutator.

Description

This operator works only on binary representation and flips each bit with a given probability \( p \in (0, 1) \). Usually it is recommended to set \( p = \frac{1}{n} \) where \( n \) is the number of bits in the representation.

Usage

mutBitflip(ind, p = 0.1)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ind</td>
<td>[binary] Binary vector, i.e., vector with elements 0 and 1 only.</td>
</tr>
<tr>
<td>p</td>
<td>[numeric(1)] Probability to flip a single bit. Default is 0.1.</td>
</tr>
</tbody>
</table>

Value

binary

References


See Also

Other mutators: mutGauss(), mutInsertion(), mutInversion(), mutJump(), mutPolynomial(), mutScramble(), mutSwap(), mutUniform()
mutGauss

Gaussian mutator.

Description

Default Gaussian mutation operator known from Evolutionary Algorithms. This mutator is applicable only for representation="float". Given an individual $x \in \mathbb{R}^l$ this mutator adds a Gaussian distributed random value to each component of $x$, i.e., $\tilde{x}_i = x_i + \sigma N(0,1)$.

Usage

mutGauss(ind, p = 1L, sdev = 0.05, lower, upper)

Arguments

- **ind** [numeric] Numeric vector / individual to mutate.
- **p** [numeric(1)] Probability of mutation for the gauss mutation operator.
- **sdev** [numeric(1)] Standard deviation of the Gauss mutation, i.e., the mutation strength.
- **lower** [numeric] Vector of minimal values for each parameter of the decision space.
- **upper** [numeric] Vector of maximal values for each parameter of the decision space.

Value

numeric

References


See Also

Other mutators: mutBitflip(), mutInsertion(), mutInversion(), mutJump(), mutPolynomial(), mutScramble(), mutSwap(), mutUniform()
**mutInsertion**  
*Insertion mutator.*

**Description**

The Insertion mutation operator selects a position random and inserts it at a random position.

**Usage**

`mutInsertion(ind)`

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>ind</td>
<td>integer</td>
</tr>
</tbody>
</table>

Permutation of integers, i.e., vector of integer values.

**Value**

integer

**See Also**

Other mutators: `mutBitflip()`, `mutGauss()`, `mutInversion()`, `mutJump()`, `mutPolynomial()`, `mutScramble()`, `mutSwap()`, `mutUniform()`

**mutInversion**  
*Inversion mutator.*

**Description**

The Inversion mutation operator selects two positions within the chromosome at random and inverts the elements inbetween.

**Usage**

`mutInversion(ind)`

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>ind</td>
<td>integer</td>
</tr>
</tbody>
</table>

Permutation of integers, i.e., vector of integer values.

**Value**

integer
mutJump

See Also

Other mutators: mutBitflip(), mutGauss(), mutInsertion(), mutJump(), mutPolynomial(), mutScramble(), mutSwap(), mutUniform()

mutJump

Jump mutator.

Description

The jump mutation operator selects two positions within the chromosome at random, say \(a\) and \(b\) with \(a < b\). Next, all elements at positions \(b - 1, b - 2, \ldots, a\) are shifted to the right by one position and finally the element at position \(b\) is assigned to position \(a\).

Usage

mutJump(ind)

Arguments

ind [integer] Permutation of integers, i.e., vector of integer values.

Value

integer

See Also

Other mutators: mutBitflip(), mutGauss(), mutInsertion(), mutInversion(), mutPolynomial(), mutScramble(), mutSwap(), mutUniform()

mutPolynomial

Polynomial mutation.

Description

Performs a polynomial mutation as used in the SMS-EMOA algorithm. Polynomial mutation tries to simulate the distribution of the offspring of binary-encoded bit flip mutations based on real-valued decision variables. Polynomial mutation favors offspring nearer to the parent.

Usage

mutPolynomial(ind, p = 0.2, eta = 10, lower, upper)
mutScramble

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ind</td>
<td>Numeric vector / individual to mutate.</td>
</tr>
<tr>
<td>p</td>
<td>Probability of mutation for each gene of an offspring. In other words, the probability that the value (allele) of a given gene will change. Default is 0.2</td>
</tr>
<tr>
<td>eta</td>
<td>Distance parameter to control the shape of the mutation distribution. Larger values generate offspring closer to the parents. Default is 10.</td>
</tr>
<tr>
<td>lower</td>
<td>Vector of minimal values for each parameter of the decision space. Must have the same length as ind.</td>
</tr>
<tr>
<td>upper</td>
<td>Vector of maximal values for each parameter of the decision space. Must have the same length as ind.</td>
</tr>
</tbody>
</table>

Value

numeric

References


See Also

Other mutators: mutBitflip(), mutGauss(), mutInsertion(), mutInversion(), mutJump(), mutScramble(), mutSwap(), mutUniform()

Description

The Scramble mutation operator selects two positions within the chromosome at random and randomly intermixes the subsequence between these positions.

Usage

mutScramble(ind)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ind</td>
<td>Permutation of integers, i.e., vector of integer values.</td>
</tr>
</tbody>
</table>
mutSwap

Value
integer

See Also
Other mutators: mutBitflip(), mutGauss(), mutInsertion(), mutInversion(), mutJump(), mutPolynomial(), mutSwap(), mutUniform()

mutSwap  Swap mutator.

Description
Choose two positions at random and swaps the genes.

Usage
mutSwap(ind)

Arguments
ind  [integer]
Permutation of integers, i.e., vector of integer values.

Value
integer

See Also
Other mutators: mutBitflip(), mutGauss(), mutInsertion(), mutInversion(), mutJump(), mutPolynomial(), mutScramble(), mutUniform()

mutUniform  Uniform mutator.

Description
This mutation operator works on real-valued genotypes only. It selects a position in the solution vector at random and replaced it with a uniformly chosen value within the box constraints of the corresponding parameter. This mutator may proof beneficial in early stages of the optimization process, since it distributes points widely within the box constraints and thus may hinder premature convergence. However, in later stages - when fine tuning is necessary, this feature is disadvantegous.
Usage

mutUniform(ind, lower, upper)

Arguments

ind [numeric]
Numeric vector / individual to mutate.

lower [numeric]
Vector of minimal values for each parameter of the decision space.

upper [numeric]
Vector of maximal values for each parameter of the decision space.

Value
numeric

See Also
Other mutators: mutBitflip(), mutGauss(), mutInsertion(), mutInversion(), mutJump(), mutPolynomial(), mutScramble(), mutSwap()

Description

This formatter function should be applied to tables where each table cell contains a $p$-value of a statistical significance test. See toLatex for an application.

Usage

niceCellFormater(cell, alpha = 0.05)

Arguments

cell [any]
Cell value. In the majority of cases this will be a numeric value.

alpha [numeric(1)]
Significance level of underlying statistical test. Default is 0.05.

Value

Formatted output.
normalize

See Also

Other EMOA performance assessment tools: approximateNadirPoint(). approximateRefPoints(). approximateRefSets(). computeDominanceRanking(). emoIndEps(). makeEMOAIndicator(). normalize(). plotDistribution(). plotFront(). plotScatter2d(). plotScatter3d(). toLatex().

normalize

Normalize approximations set(s).

Description

Normalization is done by subtracting the min.value for each dimension and dividing by the difference max.value - min.value for each dimension. Certain EMOA indicators require all elements to be strictly positive. Hence, an optional offset is added to each element which defaults to zero.

Usage

normalize(x, obj.cols, min.value = NULL, max.value = NULL, offset = NULL)

Arguments

x

[ matrix | data.frame ]

Either a numeric matrix (each column corresponds to a point) or a data.frame with columns at least obj.cols.

obj.cols

[ character(>= 2) ]

Column names of the objective functions.

min.value

[ numeric ]

Vector of minimal values of length nrow(x). Only relevant if x is a matrix. Default is the row-wise minimum of x.

max.value

[ numeric ]

Vector of maximal values of length nrow(x). Only relevant if x is a matrix. Default is the row-wise maximum of x.

offset

[ numeric ]

Numeric constant added to each normalized element. Useful to make all objectives strictly positive, e.g., located in [1, 2].

Value

[ matrix | data.frame ]

Note

In case a data.frame is passed and a “prob” column exists, normalization is performed for each unique element of the “prob” column independently (if existent).
See Also

Other EMOA performance assessment tools: approximateNadirPoint(), approximateRefPoints(), approximateRefSets(), computeDominanceRanking(), emoaiIndEps(), makeEMOAIndicator(), niceCellFormatter(), plotDistribution(), plotFront(), plotScatter2d(), plotScatter3d(), toLatex()

nsGA2

Implementation of the NSGA-II EMOA algorithm by Deb.

Description

The NSGA-II merges the current population and the generated offspring and reduces it by means of
the following procedure: It first applies the non dominated sorting algorithm to obtain the nondom-
inated fronts. Starting with the first front, it fills the new population until the i-th front does not fit.
It then applies the secondary crowding distance criterion to select the missing individuals from the
i-th front.

Usage

nsGA2(
  fitness.fun,
  n.objectives = NULL,
  n.dim = NULL,
  minimize = NULL,
  lower = NULL,
  upper = NULL,
  mu = 100L,
  lambda = mu,
  mutator = setup(mutPolynomial, eta = 25, p = 0.2, lower = lower, upper = upper),
  recombinator = setup(recSBX, eta = 15, p = 0.7, lower = lower, upper = upper),
  terminators = list(stopOnIters(100L)),
  ...
)

Arguments

  fitness.fun    [function]
                 The fitness function.

  n.objectives   [integer(1)]
                 Number of objectives of obj.fun. Optional if obj.fun is a benchmark function
                 from package smooF.

  n.dim          [integer(1)]
                 Dimension of the decision space.

  minimize       [logical(n.objectives)]
                 Logical vector with ith entry TRUE if the ith objective of fitness.fun shall
                 be minimized. If a single logical is passed, it is assumed to be valid for each
                 objective.
**lower** [numeric]
Vector of minimal values for each parameter of the decision space in case of float or permutation encoding. Optional if `obj.fun` is a benchmark function from package `smoof`.

**upper** [numeric]
Vector of maximal values for each parameter of the decision space in case of float or permutation encoding. Optional if `obj.fun` is a benchmark function from package `smoof`.

**mu** [integer(1)]
Number of individuals in the population. Default is 100.

**lambda** [integer(1)]
Offspring size, i.e., number of individuals generated by variation operators in each iteration. Default is 100.

**mutator** [ecr_mutator]
Mutation operator of type `ecr_mutator`.

**recombinator** [ecr_recombinator]
Recombination operator of type `ecr_recombinator`.

**terminators** [list]
List of stopping conditions of type “ecr_terminator”. Default is to stop after 100 iterations.

... [any]
Further arguments passed down to fitness function.

**Value**
ecr_multi_objective_result

**Note**
This is a pure R implementation of the NSGA-II algorithm. It hides the regular ecr interface and offers a more R like interface while still being quite adaptable.

**References**

---

**plotDistribution**
Plot distribution of EMOA indicators.

**Description**
Visualizes of empirical distributions of unary EMOA indicator based on the results of `computeIndicators`. 
plotDistribution

Usage

plotDistribution(
  inds,
  plot.type = "boxplot",
  fill = "algorithm",
  facet.type = "grid",
  facet.args = list(),
  logscale = character()
)

Arguments

inds  [data.frame]
  Data frame with columns “algorithm”, “prob”, “repl” and one additional column
  per EMOA indicator.

plot.type  [character(1)]
  Either “boxplot” (the default) for boxplots or “violin” for violin plots.

fill  [character(1)]
  Variable used to fill boxplots. Default is “algorithm”.

facet.type  [character(1)]
  Which faceting method to use? Pass “wrap” for facet_wrap or “grid” for
  facet_grid. Default is “wrap”.

facet.args  [list]
  Named list of arguments passed down to facet_wrap or facet_grid respectively
  (depends on facet.type). E.g., nrow to change layout. Default is the
  empty list. In this case data is grouped by problem and indicator.

logscale  [character]
  Vector of indicator names which should be log-transformed prior to visualization.
  Default is the empty character vector.

Value

ggplot

See Also

Other EMOA performance assessment tools: approximateNadirPoint(), approximateRefPoints(),
approximateRefSets(), computeDominanceRanking(), emoaIndEps(), makeEMOAIndicator(),
niceCellFormatter(), normalize(), plotFront(), plotScatter2d(), plotScatter3d(), toLatex()
**plotFront**

*Draw scatterplot of Pareto-front approximation*

**Description**

The function expects a data.frame or a matrix. By default the first 2 or 3 columns/rows are assumed to contain the elements of the approximation sets. Depending on the number of numeric columns (in case of a data.frame) or the number of rows (in case of a matrix) the function internally calls `plotScatter2d` or `plotScatter3d`.

**Usage**

```r
plotFront(x, obj.names = NULL, minimize = TRUE, ...)
```

**Arguments**

- `x`  
  [matrix|data.frame]  
  Object which contains the approximations sets.

- `obj.names`  
  [character]  
  Optional objectives names. Default is `c("f1", "f2")`.

- `minimize`  
  [logical]  
  Logical vector with ith entry TRUE if the ith objective shall be minimized. If a single logical is passed, it is assumed to be valid for each objective. If the matrix is of type `ecr_fitness_matrix` (this is the case if it is produced by one of ecr2’s utility functions, e.g. `evaluateFitness`), the appended minimize attribute is the default.

- `...`  
  [any]  
  Not used at the moment.

**Value**

`ggplot` object.

**See Also**

Other EMOA performance assessment tools: `approximateNadirPoint()`, `approximateRefPoints()`, `approximateRefSets()`, `computeDominanceRanking()`, `emoaIndEps()`, `makeEMOAIndicator()`, `niceCellFormatter()`, `normalize()`, `plotDistribution()`, `plotScatter2d()`, `plotScatter3d()`, `toLatex()`
plotHeatmap  

**Plot heatmap.**

**Description**

Given a matrix or list of matrices x this function visualizes each matrix with a heatmap.

**Usage**

```r
plotHeatmap(x, value.name = "Value", show.values = FALSE)
```

**Arguments**

- `x`  
  [matrix|list[matrix]]  
  Either a matrix or a list of matrices.

- `value.name`  
  [character(1)]  
  Name for the values represented by the matrix. Internally, the matrix is transformed into a data.frame via `melt` in order to obtain a format which may be processed by `ggplot` easily. Default is “Value”.

- `show.values`  
  [logical(1L)]  
  Should the values be printed within the heatmap cells? Default is FALSE.

**Value**

`ggplot` ggplot object.

**Examples**

```r
# simulate two (correlation) matrices
x = matrix(runif(100), ncol = 10)
y = matrix(runif(100), ncol = 10)
## Not run:
pl = plotHeatmap(x)
pl = plotHeatmap(list(x, y), value.name = "Correlation")
pl = plotHeatmap(list(MatrixX = x, MatrixY = y), value.name = "Correlation")
## End(Not run)
```
plotScatter2d

Visualize bi-objective Pareto-front approximations.

Description

Given a data frame with the results of (multiple) runs of (multiple) different multi-objective op-
timization algorithms on (multiple) problem instances the function generates `ggplot` plots of the
obtained Pareto-front approximations.

Usage

```r
plotScatter2d(  
  df,  
  obj.cols = c(“f1”, “f2”),  
  shape = “algorithm”,  
  colour = NULL,  
  highlight.algos = NULL,  
  offset.highlighted = 0,  
  title = NULL,  
  subtitle = NULL,  
  facet.type = “wrap”,  
  facet.args = list()  
)
```

Arguments

- **df** [data.frame]
  Data.frame with columns at least `obj.cols`, “prob” and “algorithm”.

- **obj.cols** [character(>= 2)]
  Column names of the objective functions. Default is `c(“f1”, “f2”).`

- **shape** [character(1)]
  Name of column which shall be used to define shape of points. Default is “algorithm”.

- **colour** [character(1)]
  Name of column which shall be used to define colour of points. Default is `NULL`,
i.e., coloring is deactivated.

- **highlight.algos** [character(1)]
  Name of algorithm to highlight exclusively. Useful to highlight, e.g., the true Pareto-optimal front (if known) or some reference set. Default is `NULL`, i.e., unknown.

- **offset.highlighted** [numeric(1)]
  Numeric offset used to shift set (see `highlight.algos`) which should be highlighted. Even though this produces objective vectors it may be used to make visible reference sets which otherwise would be hidden by overlap of multiple other approximation sets.
plotScatter2d

- **title** [character(1)]
  - Plot title.

- **subtitle** [character(1)]
  - Plot subtitle.

- **facet.type** [character(1)]
  - Which faceting method to use? Pass “wrap” for `facet_wrap` or “grid” for `facet_grid`. Default is “wrap”.

- **facet.args** [list]
  - Named list of arguments passed down to `facet_wrap` or `facet_grid` respectively (depends on `facet.type`). E.g., `nrow` to change layout. Default is the empty list. In this case data is grouped by problem.

**Value**

- **ggplot** A ggplot object.

**Note**

- At the moment only approximations of bi-objective functions are supported.

**See Also**

- Other EMOA performance assessment tools: `approximateNadirPoint()`, `approximateRefPoints()`, `approximateRefSets()`, `computeDominanceRanking()`, `emoaIndEps()`, `makeEMOAIndicator()`, `niceCellFormatter()`, `normalize()`, `plotDistribution()`, `plotFront()`, `plotScatter3d()`, `toLatex()`

**Examples**

```r
## Not run:
# load exemplary data
data(mcMST)
print(head(mcMST))

# no customization; use the defaults
pl = plotFronts(mcMST)

# algo PRIM is obtained by weighted sum scalarization
# Since the front is (mainly) convex we highlight these solutions
pl = plotFronts(mcMST, highlight.algos = "PRIM")

# customize layout
pl = plotFronts(mcMST, title = "Pareto-approximations",
                subtitle = "based on different mcMST algorithms.",
                facet.args = list(nrow = 2))

## End(Not run)
```
plotScatter3d

Visualize three-objective Pareto-front approximations.

Description

Given a data frame with the results of (multiple) runs of (multiple) different three-objective optimization algorithms on (multiple) problem instances the function generates 3D scatterplots of the obtained Pareto-front approximations.

Usage

plotScatter3d(
  df,
  obj.cols = c("f1", "f2", "f3"),
  max.in.row = 4L,
  package = "scatterplot3d",
  ...
)

Arguments

df [data.frame]  
Data.frame with columns at least obj.cols, “prob” and “algorithm”.

obj.cols [character(>= 3)]  
Column names of the objective functions. Default is c("f1", "f2", "f3").

max.in.row [integer(1)]  
Maximum number of plots to be displayed side by side in a row. Default is 4.

package [character(1L)]  
Which package to use for 3d scatterplot generation? Possible choices are “scatterplot3d”, “plot3D”, “plot3Drgl” or “plotly”. Default is “scatterplot3d”.

... [any]  
Further arguments passed down to scatterplot function.

Value

Nothing

See Also

Other EMOA performance assessment tools: approximateNadirPoint(), approximateRefPoints(), approximateRefSets(), computeDominanceRanking(), emoaIndEps(), makeEMOAIndicator(), niceCellFormatter(), normalize(), plotDistribution(), plotFront(), plotScatter2d(), toLatex()
plotStatistics  
*Generate line plot of logged statistics.*

**Description**

Expects a data.frame of logged statistics, e.g., extracted from a logger object by calling `getStatistics`, and generates a basic line plot. The plot is generated with the ggplot2 package and the ggplot object is returned.

**Usage**

```r
plotStatistics(x, drop.stats = character(0L))
```

**Arguments**

- **x**  
  [ecr_statistics|ecr_logger]  
  Logger object or statistics data frame from logger object.

- **drop.stats**  
  [character]  
  Names of logged statistics to be dropped. Default is the empty character, i.e., not to drop any stats.

---

**recCrossover**  
*One-point crossover recombinator.*

**Description**

The one-point crossover recombinator is defined for float and binary representations. Given two real-valued/binary vectors of length n, the selector samples a random position i between 1 and n-1. In the next step it creates two children. The first part of the first child contains of the subvector from position 1 to position i of the first parent, the second part from position i+1 to n is taken from the second parent. The second child is build analogously. If the parents are list of real-valued/binary vectors, the procedure described above is applied to each element of the list.

**Usage**

```r
recCrossover(inds)
```

**Arguments**

- **inds**  
  [list]  
  Parents, i.e., list of exactly two numeric or binary vectors of equal length.

**Value**

list
**recIntermediate**

Intermediate recombinator.

**Description**

Intermediate recombination computes the component-wise mean value of the \( k \) given parents. It is applicable only for float representation.

**Usage**

recIntermediate(inds)

**Arguments**

inds [list] Parents, i.e., list of exactly two numeric vectors of equal length.

**Value**

numeric Single offspring.

**See Also**

Other recombinators: recIntermediate(), recOX(), recPMX(), recSBX(), recUnifCrossover()

---

**recOX**

Ordered-Crossover (OX) recombinator.

**Description**

This recombination operator is specifically designed for permutations. The operators chooses two cut-points at random and generates two offspring as follows: a) copy the subsequence of one parent and b) remove the copied node indizes from the entire sequence of the second parent from the second cut point and b) fill the remaining gaps with this trimmed sequence.

**Usage**

recOX(inds)

**Arguments**

inds [list] Parents, i.e., list of exactly two permutations (vectors of integer values) of equal length.
**recSBX**

Value

list

See Also

Other recombinators: `recCrossover()`, `recIntermediate()`, `recPMX()`, `recSBX()`, `recUnifCrossover()`

---

**recPMX**

*Partially-Mapped-Crossover (PMX) recombinaotor.*

Description

This recombination operator is specifically designed for permutations. The operators chooses two cut-points at random and generates two offspring as follows: a) copy the subsequence of one parent and b) fill the remaining positions while preserving the order and position of as many genes as possible.

Usage

```r
recPMX(inds)
```

Arguments

- `inds` [numeric]
  Parents, i.e., list of exactly two permutations of equal length.

Value

ecr_recombinator

See Also

Other recombinators: `recCrossover()`, `recIntermediate()`, `recOX()`, `recSBX()`, `recUnifCrossover()`

---

**recSBX**

*Simulated Binary Crossover (SBX) recombinaotor.*

Description

The Simulated Binary Crossover was first proposed by [1]. It is suited for float representation only and creates two offspring. Given parents $p_1, p_2$ the offspring are generated as $c_{1/2} = \bar{x} \pm \frac{1}{2} \beta (p_2 - p_1)$ where $\bar{x} = \frac{1}{2} (p_1 + p_2), p_2 > p_1$. This way $\bar{c} = \bar{x}$ is assured.

Usage

```r
recSBX(inds, eta = 5, p = 1, lower, upper)
```
**recUnifCrossover**

**Arguments**

- **inds** [numeric]
  Parents, i.e., list of exactly two numeric vectors of equal length.
- **eta** [numeric(1)]
  Parameter eta, i.e., the distance parameters of the crossover distribution.
- **p** [numeric(1)]
  Crossover probability for each gene. Default is 1.0.
- **lower** [numeric]
  Vector of minimal values for each parameter of the decision space.
- **upper** [numeric]
  Vector of maximal values for each parameter of the decision space.

**Value**

ecr_recombinator

**Note**

This is the default recombination operator used in the NSGA-II EMOA (see nsga2).

**References**


**See Also**

Other recombinators: recCrossover(), recIntermediate(), recOX(), recPMX(), recUnifCrossover()

**Description**

Produces two child individuals. The i-th gene is from parent1 with probability p and from parent2 with probability 1−p.

**Usage**

```r
recUnifCrossover(inds, p = 0.5)
```

**Arguments**

- **inds** [list]
  Parents, i.e., list of exactly two numeric or binary vectors of equal length.
- **p** [numeric(1)]
  Probability to select gene from parent1.
**Value**

list

**See Also**

Other recombinators: recCrossover(), recIntermediate(), recOX(), recPMX(), recSBX()

---

**reduceToSingleDataFrame**

*Combine multiple data frames into a single data.frame.*

**Description**

Combine multiple data frames into a single data.frame.

**Usage**

reduceToSingleDataFrame(res = list(), what = NULL, group.col.name)

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>res</td>
<td>[list] List of data frames or other lists which contain a data frame as one of the components which is selected by what. If res is a named list those names are used to fill the group column. Otherwise the names are 1 to length(res) by default.</td>
</tr>
<tr>
<td>what</td>
<td>[character(1)] Which component of each list element in res to choose. Set this to NULL, if res is not complex, i.e., is not a list of lists.</td>
</tr>
<tr>
<td>group.col.name</td>
<td>[character(1)] Name for the grouping column.</td>
</tr>
</tbody>
</table>

---

**registerECROperator**

*Register operators to control object.*

**Description**

In ecr the control object stores information on the fitness function and serves as a storage for evolutionary components used by your evolutionary algorithm. This function handles the registration process.

**Usage**

registerECROperator(control, slot, fun, ...)


Arguments

control [ecr_control]  
Control object.

slot [character(1)]  
Name of the field in the control object where to store the operator.

fun [function]  
Actual operator. In order to use the various helper functions of ecr one needs to stick to a simple convention: The first argument of function should be the individual to mutate, a list of individuals for recombination or a matrix of fitness values for recombination. If one does not want to use the corresponding helpers, e.g., mutate, the signature of the function does not matter. However, in this case you are responsible to pass arguments correctly.

... [any]  
Further arguments for fun. These arguments are stored in the control object and passed on to fun.

Value

ecr_control

replace (mu + lambda) selection

Description

Takes a population of mu individuals and another set of lambda offspring individuals and selects mu individuals out of the union set according to the survival selection strategy stored in the control object.

Usage

replaceMuPlusLambda(
  control, population, offspring,
  fitness = NULL,
  fitness.offspring = NULL
)

replaceMuCommaLambda(
  control, population, offspring,
  fitness = NULL,
  fitness.offspring = NULL,
  n.elite = base::max(ceiling(length(population) * 0.1), 1L)
)
Arguments

control [ecr_control]
Control object.

population [list]
Current set of individuals.

offspring [list]
Another set of individuals.

fitness [matrix]
Matrix of fitness values for the individuals from population. This is only optional in the case that each individual in population has an attribute “fitness”.

fitness.offspring [matrix]
Matrix of fitness values for the individuals from offspring. This is only optional in the case that each individual in offspring has an attribute “fitness”.

n.elite [integer(1)]
Number of fittest individuals of the current generation that shall be copied to the next generation without changing. Keep in mind, that the algorithm does not care about this option if the survival.strategy is set to 'plus'. Default is 0.

Value

list List with selected population and corresponding fitness matrix.

Description

Dominated Hypervolume selector.

Performs non-dominated sorting and drops the individual from the last front with minimal hypervolume contribution. This selector is the basis of the S-Metric Selection Evolutionary Multi-Objective Algorithm, termed SMS-EMOA (see smsemoa).

Usage

selDomHV(fitness, n.select, ref.point)

Arguments

fitness [matrix]
Matrix of fitness values (each column contains the fitness value(s) of one individual).

n.select [integer(1)]
Number of elements to select.

ref.point [numeric]
Reference point for hypervolume computation.
Value

integer  Vector of survivor indizes.

Note

Note that the current implementation expects \( n.\text{select} = ncol(\text{fitness}) - 1 \) and the selection process quits with an error message if \( n.\text{select} \) is greater than 1.

See Also

Other selectors: \texttt{selGreedy()}, \texttt{selNondom()}, \texttt{selRanking()}, \texttt{selRoulette()}, \texttt{selSimple()}, \texttt{selTournament()}

Description

This utility functions expect a control object, a matrix of fitness values - each column containing the fitness value(s) of one individual - and the number of individuals to select. The corresponding selector, i.e., mating selector for \texttt{selectForMating} or survival selector for \texttt{selectForSurvival} is than called internally and a vector of indizes of selected individuals is returned.

Usage

\[
\begin{align*}
\text{selectForMating} & (\text{control}, \text{fitness}, \text{n.select}) \\
\text{selectForSurvival} & (\text{control}, \text{fitness}, \text{n.select})
\end{align*}
\]

Arguments

\[
\begin{align*}
\text{control} & \quad [\text{ecr\_control}] \\
& \quad \text{Control object.} \\
\text{fitness} & \quad [\text{matrix}] \\
& \quad \text{Matrix of fitness values (each column contains the fitness value(s) of one individual).} \\
\text{n.select} & \quad [\text{integer(1)}] \\
& \quad \text{Number of individuals to select.}
\end{align*}
\]

Details

Both functions check the optimization directions stored in the task inside the control object, i.e., whether to minimize or maximize each objective, and transparently prepare/transform the fitness matrix for the selector.

Value

integer  Integer vector with the indizes of selected individuals.
**selGreedy**

*Simple selector.*

**Description**

Sorts the individuals according to their fitness value in increasing order and selects the best ones.

**Usage**

```
selGreedy(fitness, n.select)
```

**Arguments**

- **fitness** [matrix]
  Matrix of fitness values (each column contains the fitness value(s) of one individual).
- **n.select** [integer(1)]
  Number of elements to select.

**Value**

integer  Vector of survivor indices.

**See Also**

Other selectors: `selDomHV()`, `selNondom()`, `selRanking()`, `selRoulette()`, `selSimple()`, `selTournament()`

---

**selNondom**

*Non-dominated sorting selector.*

**Description**

Applies non-dominated sorting of the objective vectors and subsequent crowding distance computation to select a subset of individuals. This is the selector used by the famous NSGA-II EMOA (see `nsga2`).

**Usage**

```
selNondom(fitness, n.select)
```

**Arguments**

- **fitness** [matrix]
  Matrix of fitness values (each column contains the fitness value(s) of one individual).
- **n.select** [integer(1)]
  Number of elements to select.
$sel\text{Ranking}$

**Value**

$set\text{OfIndividuals}$

**See Also**

Other selectors: $sel\text{DomHV}$(), $sel\text{Greedy}$(), $sel\text{Ranking}$(), $sel\text{Roulette}$(), $sel\text{Simple}$(), $sel\text{Tournament}$()

---

$sel\text{Ranking}$

*Rank Selection Operator*

**Description**

Rank-based selection preserves a constant selection pressure by sorting the population on the basis of fitness, and then allocating selection probabilities to individuals according to their rank, rather than according to their actual fitness values.

**Usage**

$sel\text{Ranking}(fitness, n.select, s = 1.5, scheme = "linear")$

**Arguments**

- **fitness** [matrix]
  - Matrix of fitness values (each column contains the fitness value(s) of one individual).
- **n.select** [integer(1)]
  - Number of elements to select.
- **s** [numeric(1)]
  - Selection pressure for linear ranking scheme with value range $[0, 1]$. Ignored if scheme is set to “exponential”. Default is 1.5.
- **scheme** [character(1)]
  - Mapping from rank number to selection probability, either “linear” or “exponential”.

**Value**

$set\text{OfIndividuals}$

**References**


**See Also**

Other selectors: $sel\text{DomHV}$(), $sel\text{Greedy}$(), $sel\text{Nondom}$(), $sel\text{Roulette}$(), $sel\text{Simple}$(), $sel\text{Tournament}$()
**selRoulette**

**Roulette-wheel / fitness-proportional selector.**

**Description**

The chance of an individual to get selected is proportional to its fitness, i.e., better individuals get a higher chance to take part in the reproduction process. Low-fitness individuals however, have a positive fitness as well.

**Usage**

selRoulette(fitness, n.select, offset = 0.1)

**Arguments**

- **fitness** [matrix]
  Matrix of fitness values (each column contains the fitness value(s) of one individual).

- **n.select** [integer(1)]
  Number of elements to select.

- **offset** [numeric(1)]
  In case of negative fitness values all values are shifted towards positive values by adding the negative of the minimal fitness value. However, in this case the minimal fitness value after the shifting process is zero. The offset is a positive numeric value which is added additionally to each shifted fitness value. This way even the individual with the smallest fitness value has a positive probability to be selected. Default is 0.1.

**Details**

Fitness proportional selection can be naturally applied to single objective maximization problems. However, negative fitness values can are problematic. The Roulette-Wheel selector thus works with the following heuristic: if negative values occur, the negative of the smallest fitness value is added to each fitness value. In this case to avoid the smallest shifted fitness value to be zero and thus have a zero probability of being selected an additional positive constant offset is added (see parameters).

**Value**

setOfIndividuals

**See Also**

Other selectors: selDomHV(), selGreedy(), selNondom(), selRanking(), selSimple(), selTournament()
**selSimple**

**Simple (naive) selector.**

**Description**

Just for testing. Actually does not really select, but instead returns a random sample of \(\text{ncol(fitness)}\) indices.

**Usage**

\[
\text{selSimple(fitness, n.select)}
\]

**Arguments**

- **fitness** [matrix]
  - Matrix of fitness values (each column contains the fitness value(s) of one individual).
- **n.select** [integer(1)]
  - Number of elements to select.

**Value**

setOfIndividuals

**See Also**

Other selectors: `selDomHV()`, `selGreedy()`, `selNondom()`, `selRanking()`, `selRoulette()`, `selTournament()`

**selTournament**

\(k\)-Tournament selector.

**Description**

\(k\) individuals from the population are chosen randomly and the best one is selected to be included into the mating pool. This process is repeated until the desired number of individuals for the mating pool is reached.

**Usage**

\[
\text{selTournament(fitness, n.select, k = 3L)}
\]
Arguments

fitness [matrix]
Matrix of fitness values (each column contains the fitness value(s) of one individual).

n.select [integer(1)]
Number of elements to select.

k [integer(1)]
Number of individuals to participate in each tournament. Default is 2L.

Value

integer Vector of survivor indizes.

See Also

Other selectors: selDomHV(), selGreedy(), selNondom(), selRanking(), selRoulette(), selSimple()

Description

The function checks, whether each points of the second set of points is dominated by at least one point from the first set.

Usage

setDominates(x, y)

Arguments

x [matrix]
First set of points.

y [matrix]
Second set of points.

Value

logical(1)
setup

Set up parameters for evolutionary operator.

Description

This function builds a simple wrapper around an evolutionary operator, i.e., mutator, recombinator or selector and defines its parameters. The result is a function that does not longer depend on the parameters. E.g., `fun = setup(mutBitflip, p = 0.3)` initializes a bitflip mutator with mutation probability 0.3. Thus, the following calls have the same behaviour: `fun(c(1, 0, 0))` and `mutBitflip(fun(c(1, 0, 0), p = 0.3))`. Basically, this type of preinitialization is only necessary if operators with additional parameters shall be initialized in order to use the black-box `ecr`.

Usage

```
setup(operator, ...)
```

Arguments

- `operator` 
  
  [ecr_operator]
  
  Evolutionary operator.

- `...` 
  
  [any]
  
  Further parameters for operator.

Value

function  Wrapper evolutionary operator with parameters x and ....

Examples

```
# initialize bitflip mutator with p = 0.3
bf = setup(mutBitflip, p = 0.3)
# sample binary string
x = sample(c(0, 1), 100, replace = TRUE)

set.seed(1)
# apply preinitialized function
print(bf(x))

set.seed(1)
# apply raw function
print(mutBitflip(x, p = 0.3))

# overwrite preinitialized values with mutate
ctrl = initECRControl(fitness.fun = function(x) sum(x), n.objectives = 1L)
# here we define a mutation probability of 0.3
ctrl = registerECROperator(ctrl, "mutate", setup(mutBitflip, p = 0.3))
# here we overwrite with 1, i.e., each bit is flipped
print(x)
print(mutate(ctrl, list(x), p.mut = 1, p = 1)[[1]])
```
Description

Default monitor object that outputs messages to the console based on a default logger (see initLogger).

Usage

setupECRDefaultMonitor(step = 10L)

Arguments

- **step**: [integer(1)]
  
  Number of steps of the EA between monitoring. Default is 10.

Value

ecr_monitor

smsemoa

Implementation of the SMS-EMOA by Emmerich et al.

Description

Pure R implementation of the SMS-EMOA. This algorithm belongs to the group of indicator based multi-objective evolutionary algorithms. In each generation, the SMS-EMOA selects two parents uniformly at, applies recombination and mutation and finally selects the best subset of individuals among all subsets by maximizing the Hypervolume indicator.

Usage

smsemoa(
  fitness.fun,
  n.objectives = NULL,
  n.dim = NULL,
  minimize = NULL,
  lower = NULL,
  upper = NULL,
  mu = 100L,
  ref.point = NULL,
  mutator = setup(mutPolynomial, eta = 25, p = 0.2, lower = lower, upper = upper),
  recombinator = setup(recSBX, eta = 15, p = 0.7, lower = lower, upper = upper),
  terminators = list(stopOnIters(100L)),
  ...)
)
Arguments

fitness.fun [function]
The fitness function.

n.objectives [integer(1)]
Number of objectives of obj.fun. Optional if obj.fun is a benchmark function from package smoof.

n.dim [integer(1)]
Dimension of the decision space.

minimize [logical(n.objectives)]
Logical vector with ith entry TRUE if the ith objective of fitness.fun shall be minimized. If a single logical is passed, it is assumed to be valid for each objective.

lower [numeric]
Vector of minimal values for each parameter of the decision space in case of float or permutation encoding. Optional if obj.fun is a benchmark function from package smoof.

upper [numeric]
Vector of maximal values for each parameter of the decision space in case of float or permutation encoding. Optional if obj.fun is a benchmark function from package smoof.

mu [integer(1)]
Number of individuals in the population. Default is 100.

ref.point [numeric]
Reference point for the hypervolume computation. Default is (11, ..., 11)' with the corresponding dimension.

mutator [ecr_mutator]
Mutation operator of type ecr_mutator.

recombinator [ecr_recombinator]
Recombination operator of type ecr_recombinator.

terminators [list]
List of stopping conditions of type “ecr_terminator”. Default is to stop after 100 iterations.

... [any]
Further arguments passed down to fitness function.

Value
ecr_multi_objective_result

Note
This helper function hides the regular ecr interface and offers a more R like interface of this state of the art EMOA.
sortByObjective

Sort Pareto-front approximation by objective.

Description

Sort Pareto-front approximation by objective.

Usage

sortByObjective(x, obj = 1L, ...)

## S3 method for class 'data.frame'
sortByObjective(x, obj = 1L, ...)

## S3 method for class 'matrix'
sortByObjective(x, obj = 1L, ...)

## S3 method for class 'ecr_multi_objective_result'
sortByObjective(x, obj = 1L, ...)

## S3 method for class 'list'
sortByObjective(x, obj = 1L, ...)

Arguments

x [object]
Object of type data frame (objectives column-wise), matrix (objectives row-wise), ecr_multi_objective_result or list (with components "pareto.front") and "pareto.set".

obj [integer(1) | character(1)]
Either the row/column number to sort by or the column name, e.g., for data frames.

... [any]
Further arguments passed down to order.

Value

Modified object.
stoppingConditions

stoppingConditions  Stopping conditions

Description

Stop the EA after a fixed number of fitness function evaluations, after a predefined number of
 generations/iterations, a given cutoff time or if the known optimal function value is approximated
 (only for single-objective optimization).

Usage

stopOnEvals(max.evals = NULL)

stopOnIters(max.iter = NULL)

stopOnOptY(opt.y, eps)

stopOnMaxTime(max.time = NULL)

Arguments

max.evals  [integer(1)]
Maximal number of function evaluations. Default is Inf.

max.iter  [integer(1)]
Maximal number of iterations/generations. Default is Inf.

opt.y  [numeric(1)]
Optimal scalar fitness function value.

eps  [numeric(1)]
Stop if absolute deviation from opt.y is lower than eps.

max.time  [integer(1)]
Time limit in seconds. Default is Inf.

Value
ecr_terminator

toGG  Transform to long format.

Description

Transform the data.frame of logged statistics from wide to ggplot2-friendly long format.
Usage

toGG(x, drop.stats = character(0L))

Arguments

x [ecr_statistics|ecr_logger]
Logger object or statistics data frame from logger object.

drop.stats [character]
Names of logged statistics to be dropped. Default is the empty character, i.e., not to drop any stats.

Value

data.frame

Description

Exports results of statistical tests to LaTeX tables.

Returns high-quality LaTeX-tables of the test results of statistical tests performed with function test on per-instance basis. i.e., a table is returned for each instances combining the results of different indicators.

Usage

toLatex(
  stats,
  stat.cols = NULL,
  probs = NULL,
  type = "by.instance",
  cell.formatter = NULL
)

## S3 method for class 'list'

## S3 method for class 'data.frame'


toParetoDf =

stat.cols = NULL,
probs = NULL,
type = "by.instance",
cell.formatter = NULL
)

Arguments

stats [list]  Data frame (return value of computeIndicators) or named list of list as returned by test.
stat.cols [character]  Names of the indicators to consider. Defaults to all indicators available in stats.
probs [character]  Filtering: vector of problem instances. This way one can restrict the size of the table(s). Defaults to all problems available in stats. Ignored if stats is a data frame.
type [character(1)]  Type of tables. At the moment only option “by.instance” is available. I.e., a separate LaTeX-table is generated for each instance specified via probs. Ignored if stats is a data frame.
cell.formatter [function(cell, ...)]  Function which is used to format table cells. This function is applied to each table cell and may be used to customize the output. Default is niceCellFormatter. Ignored if stats is a data frame.

Value

list  Named list of strings (LaTeX tables). Names correspond to the selected problem instances in probs.

See Also

Other EMOA performance assessment tools: approximateNadirPoint(), approximateRefPoints(), approximateRefSets(), computeDominanceRanking(), emoIndEps(), makeEMOAIndicator(), niceCellFormatter(), normalize(), plotDistribution(), plotFront(), plotScatter2d(), plotScatter3d()

toParetoDf  Convert matrix to Pareto front data frame.

Description

Inside ecr EMOA algorithms the fitness is maintained in an \((o, n)\) matrix where \(o\) is the number of objectives and \(n\) is the number of individuals. This function basically transposes such a matrix and converts it into a data frame.
transformFitness

Usage

```
toParetoDf(x, filter.dups = FALSE)
```

Arguments

- `x`: [matrix] Matrix.
- `filter.dups`: [logical(1)] Shall duplicates be removed? Default is FALSE.

Value
data.frame

transformFitness  

Fitness transformation / scaling.

Description

Some selectors support maximization only, e.g., roulette wheel selector, or minimization (most others). This function computes a factor from -1, 1 for each objective to match supported selector optimization directions and the actual objectives of the task.

Usage

```
transformFitness(fitness, task, selector)
```

Arguments

- `fitness`: [matrix] Matrix of fitness values with the fitness vector of individual i in the i-th column.
- `task`: [ecr_optimization_task] Optimization task.
- `selector`: [ecr_selector] Selector object.

Value

matrix  Transformed / scaled fitness matrix.
updateLogger

Update the log.

Description

This function modifies the log in-place, i.e., without making copies.

Usage

updateLogger(log, population, fitness = NULL, n.evals, extras = NULL, ...)

Arguments

log [ecr_logger]
The log generated by initLogger.

population [list]
List of individuals.

fitness [matrix]
Optional matrix of fitness values (each column contains the fitness value(s) for one individual) of population. If no matrix is passed and the log shall store information of the fitness, each individual needs to have an attribute fitness.

n.evals [integer(1)]
Number of fitness function evaluations performed in the last generation.

extras [list]
Optional named list of additional scalar values to log. See log.extras argument of initLogger for details.

...
[any]
Further arguments. Not used at the moment.

See Also

Other logging: getPopulationFitness(), getPopulations(), getStatistics(), initLogger()

updateParetoArchive

Update Pareto Archive.

Description

This function updates a Pareto archive, i.e., an archive of non-dominated points. It expects the archive, a set of individuals, a matrix of fitness values (each column corresponds to the fitness vector of one individual) and updates the archive “in-place”. If the archive has unlimited capacity all non-dominated points of the union of archive and passed individuals are stored. Otherwise, i.e., in case the archive is limited in capacity (argument max.size of initParetoArchive was set to an integer value greater zero), the trunc.fun function passed to initParetoArchive is applied to all non-dominated points to determine which points should be dropped.
which.dominated

Usage

```r
updateParetoArchive(archive, inds, fitness, ...)
```

Arguments

- **archive**
  - `[ecr_pareto_archive]`
  - The archive generated by `initParetoArchive`.

- **inds**
  - `[list]`
  - List of individuals.

- **fitness**
  - `[matrix]`
  - Matrix of fitness values (each column contains the fitness value(s) for one individual) of `inds`.

- **...**
  - `[any]`
  - Further arguments passed down to `trunc.fun` (set via `initParetoArchive`).

See Also

- Other ParetoArchive: `getIndividuals()`, `getSize()`, `initParetoArchive()`

---

**which.dominated**

*Determine which points of a set are (non)dominated.*

Description

Given a matrix with one point per column which.dominated returns the column numbers of the dominated points and which.nondominated the column numbers of the nondominated points. Function `isMaximallyDominated` returns a logical vector with TRUE for each point which is located on the last non-domination level.

Usage

```r
which.dominated(x)

which.nondominated(x)

isMaximallyDominated(x)
```

Arguments

- **x**
  - `[matrix]`
  - Numeric (n x d) matrix where n is the number of points and d is the number of objectives.

Value

integer
Examples

```r
data(mtcars)
# assume we want to maximize horsepower and minimize gas consumption
cars = mtcars[, c("mpg", "hp")]
cars$hp = -cars$hp
idxs = which.nondominated(as.matrix(cars))
print(mtcars[idxs, ])
```

wrapChildren  

Wrap the individuals constructed by a recombination operator.

Description

Should be used if the recombinator returns multiple children.

Usage

```r
wrapChildren(...)```

Arguments

```r
... [any]
    Individuals.
```

Value

*list*  List of individuals.
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