# Package 'ExpDE'

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 $check\_stop\_criteria$  Stop criteria for DE

### Description

Implements different stop criteria for the ExpDE framework

### Usage

check\_stop\_criteria()

### Value

logical flag indicating whether any stop condition has been reached.

### Warning

This routine accesses the parent environment used in the main function ExpDE(), which means that changes made in the variables contained in env WILL change the original values. DO NOT change anything unless you're absolutely sure of what you're doing.

create\_population 3

### Description

Creates a new population for the ExpDE framework

#### Usage

```
create_population(popsize, probpars)
```

### **Arguments**

popsize population size

probpars list of named problem parameters (see ExpDE).

### Value

A matrix containing the population for the ExpDE

### **Description**

Evaluates the DE population on a given objective function.

### Usage

```
evaluate_population(probpars, Pop)
```

### Arguments

probpars problem parameters (see ExpDE for details).

Pop population matrix (each row is a candidate solution, normalized to the [0, 1]

interval,)

### Value

numeric vector (with length nrow(Pop)) containing the function values of each point in the population.

ExpDE	Experimental Differential Evolution - ExpDE

### Description

Modular implementation of the Differential Evolution Algorithm for the experimental investigation of the effects of different operators on the performance of the algorithm.

### Usage

```
ExpDE(popsize, mutpars = list(name = "mutation_rand", f = 0.2),
  recpars = list(name = "recombination_bin", cr = 0.8, nvecs = 1),
  selpars = list(name = "standard"), stopcrit, probpars, seed = NULL,
  showpars = list(show.iters = "none"))
```

### **Arguments**

popsize	population size
mutpars	list of named mutation parameters. See Mutation parameters for details.
recpars	list of named recombination parameters. See Recombination parameters for details.
selpars	list of named selection parameters. See Selection parameters for details.
stopcrit	list of named stop criteria parameters. See Stop criteria for details.
probpars	list of named problem parameters. See Problem Description for details.
seed	seed for the random number generator. See Random Seed for details.
showpars	parameters that regulate the echoing of progress indicators See Showpars for details.

### **Details**

This routine is used to launch a differential evolution algorithm for the **minimization** of a given problem instance using different variants of the recombination, mutation and selection operators. The input parameters that describe those operators receive list objects describing the operator variants to be used in a given optimization procedure.

#### Value

A list object containing the final population (sorted by performance) , the performance vector, and some run statistics.

#### **Mutation Parameters**

mutpars is used to inform the routine the type of differential mutation to use, as well as any mutation-related parameter values. The current version accepts the following options:

- mutation\_best
- mutation\_rand
- mutation\_mean
- mutation\_none
- mutation\_current\_to\_pbest (incl. special case current-to-best)
- mutation\_wgi

mutpars receives a list object with name field mutparsname (containing the name of the function to be called, e.g., name = "mutation\_rand") as well as whatever parameters that function may require/accept (e.g., mutparsf = 0.7, mutparsnvecs = 2, etc.). See the specific documentation of each function for details.

Some examples are provided in the Examples section below.

#### **Recombination parameters**

As with the mutation parameters, recpars is used to define the desired recombination strategy. The current version accepts the following options:

- recombination\_arith
- recombination\_bin
- recombination\_blxAlphaBeta (incl. special cases blxAlpha and flat)
- recombination\_eigen
- recombination\_exp
- recombination\_geo
- recombination\_lbga
- recombination\_linear
- recombination\_mmax
- recombination\_npoint
- recombination\_none
- recombination\_onepoint
- recombination\_pbest
- recombination\_sbx
- recombination\_wright

recpars receives a list object with name field recpars\$name (containing the name of the function to be called, e.g., name = "recombination\_bin") as well as whatever parameters that function may require/accept (e.g., recpars\$cr = 0.8, recpars\$minchange = TRUE, etc.). See the specific documentation of each function for details.

Some examples are provided in the Examples section below.

#### **Selection parameters**

selpars follows the same idea as mutpars and recpars, and is used to define the selection operators. Currently, only the standard DE selection, selection\_standard, is implemented.

### Stop criteria

stopcrit is similar to recpar and the other list arguments, but with the difference that multiple stop criteria can be defined for the algorithm. The names of the stop criteria to be used are passed in the stopcrit\$names field, which must contain a character vector. Other parameters to be used for stopping the algorithm (e.g., the maximum number of iterations stopcrit\$maxiter) can also be included as stopcrit fields. Currently implemented criteria are:

- "stop\_maxiter" (requires additional field stopcrit\$maxiter = ? with the maximum number of iterations).
- "stop\_maxeval" (requires additional field stopcrit\$maxevals = ? with the maximum number of function calls).

See check\_stop\_criteria for details.

### **Problem description**

The probpers parameter receives a list with all definitions related to the problem instance to be optimized. There are three required fields in this parameter:

- probpars\$name, the name of the function that represents the problem to be solved.
- probpars\$xmin, a vector containing the lower bounds of all optimization variables (i.e., a vector of length M, where M is the dimension of the problem).
- probpars\$xmax, a vector containing the upper bounds of all optimization variables.

This list can also contain the following optional arguments

• probpars\$matrixEval, indicates what kind of input is expected by the function provided in probpars\$name. Valid entries are "vector", "colMatrix" and "rowMatrix". Defaults to probpars\$matrixEval = "rowMatrix"

**Important**: the objective function routine must receive either a vector or a matrix of vectors to be evaluated in the form of an input parameter named either "x" or "X" or "Pop" (any one of the three is allowed).

#### Random Seed

The seed argument receives the desired seed for the PRNG. This value can be set for reproducibility purposes. The value of this parameter defaults to NULL, in which case the seed is arbitrarily set using as.numeric(Sys.time()).

#### **Showpars**

showpars is a list containing parameters that control the printed output of ExpDE. Parameter showpars can have the following fields:

- showpars\$show.iters = c("dots", "numbers", "none"): type of output. Defaults to "numbers".
- showpars\$showevery: positive integer that determines how frequently the routine echoes something to the terminal. Defaults to 1.

### Author(s)

Felipe Campelo (<fcampelo@ufmg.br>) and Moises Botelho (<moisesufop@gmail.com>)

#### References

F. Campelo, M. Botelho, "Experimental Investigation of Recombination Operators for Differential Evolution", Genetic and Evolutionary Computation Conference, July 20-24, 2016, Denver/CO. DOI: 10.1145/2908812.2908852

### **Examples**

```
# DE/rand/1/bin with population 40, F = 0.8 and CR = 0.5
popsize <- 100
mutpars <- list(name = "mutation_rand", f = 0.8)</pre>
recpars <- list(name = "recombination_bin", cr = 0.5, minchange = TRUE)</pre>
selpars <- list(name = "selection_standard")</pre>
stopcrit <- list(names = "stop_maxiter", maxiter = 100)</pre>
probpars <- list(name = "sphere",</pre>
                 xmin = rep(-5.12,10), xmax = rep(5.12,10))
seed <- NULL
showpars <- list(show.iters = "numbers", showevery = 1)</pre>
ExpDE(popsize, mutpars, recpars, selpars, stopcrit, probpars, seed, showpars)
# DE/wgi/1/blxAlpha
recpars <- list(name = "recombination_blxAlphaBeta", alpha = 0.1, beta = 0.1)
mutpars <- list(name = "mutation_wgi", f = 0.8)</pre>
ExpDE(popsize, mutpars, recpars, selpars, stopcrit, probpars)
# DE/best/1/sbx
recpars <- list(name = "recombination_sbx", eta = 10)</pre>
mutpars <- list(name = "mutation_best", f = 0.6, nvecs = 1)</pre>
ExpDE(popsize, mutpars, recpars, selpars, stopcrit, probpars)
# DE/best/1/eigen/bin
recpars <- list(name = "recombination_eigen",</pre>
                  othername = "recombination_bin",
                  cr = 0.5, minchange = TRUE)
showpars <- list(show.iters = "dots", showevery = 10)</pre>
stopcrit <- list(names = "stop_maxeval", maxevals = 10000)</pre>
ExpDE(popsize, mutpars, recpars, selpars, stopcrit, probpars, seed = 1234)
```

8 mutation\_best

mutation\_best

/best mutation for DE

### **Description**

Implements the "/best/nvecs" mutation for the ExpDE framework

### Usage

```
mutation_best(X, mutpars)
```

### Arguments

X population matrix

mutpars mutation parameters (see Mutation parameters for details)

#### Value

Matrix M containing the mutated population

#### **Mutation Parameters**

The mutpars parameter contains all parameters required to define the mutation. mutation\_best() understands the following fields in mutpars:

- f : scaling factor for difference vector(s).

  Accepts numeric vectors of size 1 or nvecs.
- nvecs: number of difference vectors to use.
   Accepts 1 <= nvecs <= (nrow(X)/2 2)</li>
   Defaults to 1.

### Warning

This routine will search for the performance vector of population X (J) in the parent environment (using parent.frame(). This variable must be defined for mutation\_best() to work.

### References

K. Price, R.M. Storn, J.A. Lampinen, "Differential Evolution: A Practical Approach to Global Optimization", Springer 2005

### Author(s)

Felipe Campelo (<fcampelo@ufmg.br>)

mutation\_current\_to\_pbest

/current-to-pbest mutation for DE

### Description

Implements the "/current-to-pbest" mutation for the ExpDE framework

#### **Usage**

```
mutation_current_to_pbest(X, mutpars)
```

### **Arguments**

X population matrix

mutation parameters (see Mutation parameters for details)

#### **Details**

This routine also implements one special case:

- current-to-best mutation (current\_to\_best), by setting mutpars\$p = 1);
- Flat recombination (flat), by setting recpars\$alpha = recpars\$beta = 0)

#### Value

Matrix M containing the mutated population

### **Mutation Parameters**

The mutpars parameter contains all parameters required to define the mutation. mutation\_current\_to\_pbest() understands the following fields in mutpars:

- f : scaling factor for difference vector(s).

  Accepts numeric vectors of size 1 or nvecs.
- p: either the number of "best" vectors to use (if given as a positive integer) or the proportion of the population to use as "best" vectors (if 0 ).

### Warning

This routine will search for the performance vector of population X (J) in the parent environment (using parent.frame(). This variable must be defined for mutation\_current\_to\_pbest() to work.

#### References

J. Zhang, A.C. Sanderson, "JADE: Adaptive differential evolution with optional external archive". IEEE Transactions on Evolutionary Computation 13:945-958, 2009

10 mutation\_mean

#### Author(s)

Felipe Campelo (<fcampelo@ufmg.br>)

mutation\_mean

/mean mutation for DE

### Description

Implements the "/mean/nvecs" mutation for the ExpDE framework

### Usage

```
mutation_mean(X, mutpars)
```

### **Arguments**

X population matrix

mutpars mutation parameters (see Mutation parameters for details)

### Value

Matrix M containing the mutated population

### **Mutation Parameters**

The mutpars parameter contains all parameters required to define the mutation. mutation\_mean() understands the following fields in mutpars:

- f : scaling factor for difference vector(s).

  Accepts numeric vectors of size 1 or nvecs.
- nvecs: number of difference vectors to use.
   Accepts 1 <= nvecs <= (nrow(X)/2 2)</li>
   Defaults to 1.

#### References

K. Price, R.M. Storn, J.A. Lampinen, "Differential Evolution: A Practical Approach to Global Optimization", Springer 2005

#### Author(s)

Felipe Campelo (<fcampelo@ufmg.br>)

mutation\_none 11

mutation\_none

NULL mutation for DE

### **Description**

Implements the "/none" mutation (i.e., no mutation performed) for the ExpDE framework

#### Usage

```
mutation_none(X, mutpars)
```

### **Arguments**

X population matrix

mutpars mutation parameters (see Mutation parameters for details)

#### Value

@return The same matrix X used as an input.

### **Mutation Parameters**

The mutpars parameter contains all parameters required to define the mutation. mutation\_none() requires no fields in this parameter.

mutation\_operators

Mutation operators available

### Description

List all available mutation operators in the ExpDE package

### Usage

```
mutation_operators()
```

### Value

Character vector with the names of all mutation operators

12 mutation\_rand

mutation\_rand

/rand mutation for DE

### Description

Implements the "/rand/nvecs" mutation for the ExpDE framework

### Usage

```
mutation_rand(X, mutpars)
```

### Arguments

X population matrix

mutation parameters (see Mutation parameters for details)

### Value

Matrix M containing the mutated population

### **Mutation Parameters**

The mutpars parameter contains all parameters required to define the mutation. mutation\_rand() understands the following fields in mutpars:

- f : scaling factor for difference vector(s).

  Accepts numeric vectors of size 1 or nvecs.
- nvecs: number of difference vectors to use.
   Accepts 1 <= nvecs <= (nrow(X)/2 2)</li>
   Defaults to 1.

### References

K. Price, R.M. Storn, J.A. Lampinen, "Differential Evolution: A Practical Approach to Global Optimization", Springer 2005

#### Author(s)

Felipe Campelo (<fcampelo@ufmg.br>)

mutation\_wgi 13

mutation\_wgi

/wgi mutation for DE

### **Description**

Implements the "/wgi/nvecs" mutation (weighted global intermediate) for the ExpDE framework. This variant is based on a recombination strategy known as "weighted global intermediate recombination" (see the References section for details)

### Usage

```
mutation_wgi(X, mutpars)
```

### **Arguments**

X population matrix

mutation parameters (see Mutation parameters for details)

### Value

Matrix M containing the mutated population

### **Mutation Parameters**

The mutpars parameter contains all parameters required to define the mutation. mutation\_wgi() understands the following fields in mutpars:

- f : scaling factor for difference vector(s).

  Accepts numeric vectors of size 1 or nvecs.
- nvecs: number of difference vectors to use.
   Accepts 1 <= nvecs <= (nrow(X)/2 2)</li>
   Defaults to 1.

### References

D. Arnold, "Weighted multirecombination evolution strategies". Theoretical Computer Science 361(1): 18-37, 2006.

T. Glasmachers, C. Igel, "Uncertainty handling in model selection for support vector machines". Proc. International Conference on Parallel Problem Solving from Nature (PPSN'08), 185-194, 2008.

#### Author(s)

Felipe Campelo (<fcampelo@ufmg.br>)

14 recombination\_arith

print\_progress

Print progress of DE

### **Description**

Echoes the progress of DE to the terminal

### Usage

```
print_progress()
```

#### **Parameters**

This routine accesses all variables defined in the calling environment using parent.frame(), so it does not require any explicit input parameters. However, the calling environment must contain:

- showpars: list containing parameters that control the printed output of moead(). Parameter showpars can have the following fields:
  - \$show.iters = c("dots", "numbers", "none"): type of output. Defaults to "numbers".
  - \$showevery: positive integer that determines how frequently the routine echoes something to the terminal. Defaults to 1.
- iters(): counter function that registers the iteration number

recombination\_arith

Arithmetic recombination for DE

### **Description**

Implements the "/arith" (arithmetic) recombination for the ExpDE framework

### Usage

```
recombination_arith(X, M, ...)
```

### **Arguments**

X population matrix (original)M population matrix (mutated)... optional parameters (unused)

### Value

Matrix U containing the recombined population

recombination\_bin 15

#### References

F. Herrera, M. Lozano, A. M. Sanchez, "A taxonomy for the crossover operator for real-coded genetic algorithms: an experimental study", International Journal of Intelligent Systems 18(3) 309-338, 2003.

recombination\_bin

/bin recombination for DE

#### **Description**

Implements the "/bin" (binomial) recombination for the ExpDE framework

### Usage

```
recombination_bin(X, M, recpars)
```

### **Arguments**

X population matrix (original)
M population matrix (mutated)

recpars recombination parameters (see Recombination parameters for details)

### Value

Matrix U containing the recombined population

### **Recombination Parameters**

The recpars parameter contains all parameters required to define the recombination. recombination\_bin() understands the following fields in recpars:

- cr : component-wise probability of using the value in M.
   Accepts numeric value 0 < cr <= 1.</li>
- minchange: logical flag to force each new candidate solution to inherit at least one component from its mutated 'parent'.
   Defaults to TRUE

### References

K. Price, R.M. Storn, J.A. Lampinen, "Differential Evolution: A Practical Approach to Global Optimization", Springer 2005

recombination\_blxAlphaBeta

Blend Alpha Beta recombination for DE

### **Description**

Implements the "/blxAlphaBeta" (Blend Alpha Beta) recombination for the ExpDE framework

### Usage

```
recombination_blxAlphaBeta(X, M, recpars)
```

#### **Arguments**

X population matrix (original)
M population matrix (mutated)

recombination parameters (see Recombination parameters for details)

#### **Details**

This routine also implements two special cases:

- BLX-alpha recombination (blxAlpha), by setting recpars\$alpha = recpars\$beta);
- Flat recombination (flat), by setting recpars\$alpha = recpars\$beta = 0)

#### Value

Matrix U containing the recombined population

#### **Recombination Parameters**

The recpars parameter contains all parameters required to define the recombination. recombination\_blxAlpha() understands the following fields in recpars:

- alpha: extrapolation parameter for 'best' parent vector.
   Accepts real value 0 <= alpha <= 0.5.</li>
- beta: extrapolation parameter for 'worst' parent vector.
   Accepts real value 0 <= beta <= 0.5.</li>

@section Warning: This recombination operator evaluates the candidate solutions in M, which adds an extra popsize evaluations per iteration.

### References

F. Herrera, M. Lozano, A. M. Sanchez, "A taxonomy for the crossover operator for real-coded genetic algorithms: an experimental study", International Journal of Intelligent Systems 18(3) 309-338, 2003.

recombination\_eigen 17

recombination\_eigen /eigen recombination for DE

### **Description**

Implements the "/eigen" (eigenvector-based) recombination for the ExpDE framework

### Usage

```
recombination_eigen(X, M, recpars)
```

#### **Arguments**

X population matrix (original)M population matrix (mutated)

recpars recombination parameters (see Recombination parameters for details)

#### Value

Matrix U containing the recombined population

#### **Recombination Parameters**

The recpars parameter contains all parameters required to define the recombination. recombination\_eigen() understands the following fields in recpars:

- othername: name of the recombination operator to be applied after the projection in the eigenvector basis
- ...: parameters required (or optional) to the operator defined by recpars\$othername

#### References

Shu-Mei Guo e Chin-Chang Yang, "Enhancing differential evolution utilizing eigenvector-based crossover operator", IEEE Transactions on Evolutionary Computation 19(1):31-49, 2015.

 ${\it recombination\_exp}$   ${\it Exponential\ recombination\ for\ DE}$ 

#### **Description**

Implements the "/exp" (exponential) recombination for the ExpDE framework

### Usage

```
recombination_exp(X, M, recpars)
```

18 recombination\_geo

#### **Arguments**

X population matrix (original)
M population matrix (mutated)

recpars recombination parameters (see Recombination parameters for details)

#### Value

Matrix U containing the recombined population

#### **Recombination Parameters**

The recpars parameter contains all parameters required to define the recombination. recombination\_exp() understands the following fields in recpars:

cr : component-wise probability of selection as a cut-point.
 Accepts numeric value 0 < cr <= 1.</li>

#### References

K. Price, R.M. Storn, J.A. Lampinen, "Differential Evolution: A Practical Approach to Global Optimization", Springer 2005

recombination\_geo

Geometric recombination for DE

### **Description**

Implements the "/geo" (geometric) recombination for the ExpDE framework

### Usage

```
recombination_geo(X, M, recpars)
```

### **Arguments**

X population matrix (original)M population matrix (mutated)

recpars recombination parameters (see Recombination parameters for details)

#### Value

Matrix U containing the recombined population

recombination\_lbga 19

#### **Recombination Parameters**

The recpars parameter contains all parameters required to define the recombination. recombination\_geo() understands the following fields in recpars:

alpha: exponent for geometrical recombination.
 Accepts numeric value 0 <= alpha <= 1 or NULL (in which case a random value is chosen for each recombination).</li>

#### References

F. Herrera, M. Lozano, A. M. Sanchez, "A taxonomy for the crossover operator for real-coded genetic algorithms: an experimental study", International Journal of Intelligent Systems 18(3) 309-338, 2003.

recombination\_lbga

Linear BGA recombination for DE

### **Description**

Implements the "/lbga" (Linear Breeder Genetic Algorithm) recombination for the ExpDE framework

#### Usage

```
recombination_lbga(X, M, ...)
```

#### **Arguments**

X population matrix (original)M population matrix (mutated)... optional parameters (unused)

### Value

Matrix U containing the recombined population

### Warning

This recombination operator evaluates the candidate solutions in M, which adds an extra popsize evaluations per iteration.

#### References

- F. Herrera, M. Lozano, A. M. Sanchez, "A taxonomy for the crossover operator for real-coded genetic algorithms: an experimental study", International Journal of Intelligent Systems 18(3) 309-338, 2003.
- D. Schlierkamp-voosen, H. Muhlenbein, "Strategy Adaptation by Competing Subpopulations", Proc. Parallel Problem Solving from Nature (PPSN III), 199-208, 1994.

20 recombination\_mmax

recombination\_linear Linear recombination for DE

### **Description**

Implements the "/linear" recombination for the ExpDE framework

#### Usage

```
recombination_linear(X, M, ...)
```

### **Arguments**

X population matrix (original)
M population matrix (mutated)
... optional parameters (unused)

#### Value

Matrix U containing the recombined population

### Warning

This recombination operator evaluates 3\*popsize candidate solutions per iteration of the algorithm. The value of the nfe counter and the vector of performance values G are updated in the calling environment.

### References

F. Herrera, M. Lozano, A. M. Sanchez, "A taxonomy for the crossover operator for real-coded genetic algorithms: an experimental study", International Journal of Intelligent Systems 18(3) 309-338, 2003.

A.H. Wright, "Genetic Algorithms for Real Parameter Optimization", Proc. Foundations of Genetic Algorithms, 205-218, 1991.

recombination\_mmax

Min Max Arithmetical recombination for DE

#### **Description**

Implements the "/mmax" (min-max-arithmetical) recombination for the ExpDE framework

### Usage

```
recombination_mmax(X, M, recpars = list(lambda = NULL))
```

recombination\_none 21

#### **Arguments**

X population matrix (original)
M population matrix (mutated)

recpars recombination parameters (see Recombination parameters for details)

#### Value

Matrix U containing the recombined population

### Warning

This recombination operator evaluates 4\*popsize candidate solutions per iteration of the algorithm. The value of the nfe counter and the vector of performance values G are updated in the calling environment.

#### **Recombination Parameters**

The recpars parameter contains all parameters required to define the recombination. recombination\_pbest() understands the following fields in recpars:

• lambda: Recombination multiplier.

Optional. Defaults to NULL Accepts numeric value 0 < lambda < 1 or NULL (in which case a random value is independently used for each variable of each recombination pair).

#### References

F. Herrera, M. Lozano, A. M. Sanchez, "A taxonomy for the crossover operator for real-coded genetic algorithms: an experimental study", International Journal of Intelligent Systems 18(3):309-338, 2003

F Herrera, M. Lozano, J.L. Verdegay, "Tuning fuzzy logic controllers by genetic algorithms.", International Journal of Approximate Reasoning 12(3):299-315, 1995.

recombination\_none

NULL recombination for DE

#### **Description**

Implements the "/none" recombination (i.e., no recombination performed) for the ExpDE framework

### Usage

```
recombination_none(X, M, ...)
```

22 recombination\_npoint

### Arguments

Χ	population matrix (original)
М	population matrix (mutated)
	optional parameters (unused)

#### Value

The same matrix M used as an input.

recombination\_npoint n-point recombination for DE

### Description

Implements the "/npoint" (n-point) recombination for the ExpDE (as used in the Simple GA).

### Usage

```
recombination_npoint(X, M, recpars = list(N = NULL))
```

### **Arguments**

X population matrix (original)
M population matrix (mutated)

recpars recombination parameters (see Recombination parameters for details)

#### Value

Matrix U containing the recombined population

#### **Recombination Parameters**

The recpars parameter contains all parameters required to define the recombination. recombination\_npoint() understands the following fields in recpars:

N: cut number points for crossover.
 Accepts integer value 0 <= N < n, where n is the dimension of the problem; Use N = 0 or N = NULL for randomly choosing a number of cut points.</li>
 Defaults to NULL.

#### References

L.J. Eshelman, R.A. Caruana, J.D. Schaffer (1989), "Biases in the crossover landscape. In: Proceedings of the Third International Conference on Genetic Algorithms, pp. 10-19, San Francisco, CA, USA.

recombination\_onepoint

One-point recombination for DE

### **Description**

Implements the one-point recombination (as used in the Simple GA).

### Usage

```
recombination_onepoint(X, M, recpars = list(K = NULL))
```

### **Arguments**

X population matrix (original)

M population matrix (mutated)

recpars recombination parameters (see Recombination parameters for details)

#### Value

Matrix U containing the recombined population

#### **Recombination Parameters**

The recpars parameter contains all parameters required to define the recombination. recombination\_onepoint() understands the following fields in recpars:

K: cut point for crossover.
 Accepts integer value 0 <= K < n, where n is the dimension of the problem; Use K = 0 or K = NULL for randomly choosing a position for each pair of points.</li>
 Defaults to NULL.

#### References

F. Herrera, M. Lozano, A. M. Sanchez, "A taxonomy for the crossover operator for real-coded genetic algorithms: an experimental study", International Journal of Intelligent Systems 18(3) 309-338, 2003.

24 recombination\_pbest

recombination\_operators

Recombination operators available

### **Description**

List all available recombination operators in the ExpDE package

#### Usage

```
recombination_operators()
```

#### Value

Character vector with the names of all recombination operator routines

p-Best recombination\_pbest p-Best recombination for DE

### **Description**

Implements the "/pbest" (p-Best) recombination for the ExpDE framework

### Usage

```
recombination_pbest(X, M, recpars)
```

### **Arguments**

X population matrix (original)M population matrix (mutated)

recpars recombination parameters (see Recombination parameters for details)

#### Value

Matrix U containing the recombined population

#### **Recombination Parameters**

The recpars parameter contains all parameters required to define the recombination. recombination\_pbest() understands the following fields in recpars:

cr : component-wise probability of using the value in M.
 Accepts numeric value 0 < cr <= 1.</li>

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#### Warning

This routine will search for the iterations counter (t), the maximum number of iterations (stopcrit\$maxiter), and the performance vector of population X (J) in the parent environment (using parent.frame(). These variables must be defined for recombination\_pbest() to work.

#### References

S.M. Islam, S. Das, S. Ghosh, S. Roy, P.N. Suganthan, "An Adaptive Differential Evolution Algorithm With Novel Mutation and Crossover Strategies for Global Numerical Optimization", IEEE. Trans. Systems, Man and Cybernetics - Part B 42(2), 482-500, 2012

recombination\_sbx

/sbx recombination for DE

#### **Description**

Implements the "/sbx" (Simulated Binary) recombination for the ExpDE framework

### Usage

```
recombination_sbx(X, M, recpars)
```

### **Arguments**

X population matrix (original)
M population matrix (mutated)

recpars recombination parameters (see Recombination parameters for details)

#### Value

Matrix U containing the recombined population

#### **Recombination Parameters**

The recpars parameter contains all parameters required to define the recombination. recombination\_sbx() understands the following field in recpars:

eta: spread factor.
 Accepts numeric value eta > 0.

#### References

K. Price, R.M. Storn, J.A. Lampinen, "Differential Evolution: A Practical Approach to Global Optimization", Springer 2005

F. Herrera, M. Lozano, A. M. Sanchez, "A taxonomy for the crossover operator for real-coded genetic algorithms: an experimental study", International Journal of Intelligent Systems 18(3) 309-338, 2003.

K. Deb, R.B. Agrawal, "Simulated binary crossover for continuous search space", Complex Systems (9):115-148, 1995.

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recombination\_wright Heuristic Wright recombination for DE

### **Description**

Implements the "/wright" (Heuristic Wright) recombination for the ExpDE framework.

### Usage

```
recombination_wright(X, M, ...)
```

### Arguments

X population matrix (original)M population matrix (mutated)... optional parameters (unused)

#### Value

Matrix U containing the recombined population

#### Warning

This recombination operator evaluates the candidate solutions in M, which adds an extra popsize evaluations per iteration.

#### References

F. Herrera, M. Lozano, A. M. Sanchez, "A taxonomy for the crossover operator for real-coded genetic algorithms: an experimental study", International Journal of Intelligent Systems 18(3) 309-338, 2003.

A.H. Wright, "Genetic Algorithms for Real Parameter Optimization", Proc. Foundations of Genetic Algorithms, 205-218, 1991.

selection\_standard Standard selection for DE

### **Description**

Implements the standard selection (greedy) for the ExpDE framework

### Usage

```
selection_standard(X, U, J, G)
```

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### Arguments

X	population matrix (original)
U	population matrix (recombined)
J	performance vector for population X
G	performance vector for population U

### Value

list object containing the selected population (Xsel) and its corresponding performance values (Jsel).

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