

# Package: genalg (via r-universe)

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**Title** R Based Genetic Algorithm

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**Description** R based genetic algorithm for binary and floating point chromosomes.

**License** GPL-2

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## Description

Plots features of the genetic algorithm optimization run. The default plot shows the minimal and mean evaluation value, indicating how far the GA has progressed.

The "hist" plot shows for binary chromosome the gene selection frequency, i.e. the times one gene in the chromosome was selected in the current population. In case of floats chromosomes, it will make histograms for each variable to indicate the selected values in the population.

The "vars" plot the evaluation function versus the variable value. This is useful to look at correlations between the variable and the evaluation values.

## Usage

```
## S3 method for class 'rbga'  
plot(x, type="default", breaks=10, ...)
```

## Arguments

x	a rbga object.
type	one of "hist", "vars" or "default".
breaks	the number of breaks in a histogram.
...	options directly passed to the plot function.

## Examples

```
evaluate <- function(string=c()) {  
  returnVal = 1 / sum(string);  
  returnVal  
}  
  
rbga.results = rbga.bin(size=10, mutationChance=0.01, zeroToOneRatio=0.5,  
  evalFunc=evaluate)  
  
plot(rbga.results)  
plot(rbga.results, type="hist")
```

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 rbga

*R Based Genetic Algorithm (floating point chromosome)*


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### Description

A R based genetic algorithm that optimizes, using a user set evaluation function, a set of floats. It takes as input minimum and maximum values for the floats to optimize. The optimum is the chromosome for which the evaluation value is minimal.

It requires a `evalFunc` method to be supplied that takes as argument the chromosome, a vector of floats. Additionally, the GA optimization can be monitored by setting a `monitorFunc` that takes a `rbga` object as argument.

Results can be visualized with `plot.rbga` and summarized with `summary.rbga`.

### Usage

```
rbga(stringMin=c(), stringMax=c(),
     suggestions=NULL,
     popSize=200, iters=100,
     mutationChance=NA,
     elitism=NA,
     monitorFunc=NULL, evalFunc=NULL,
     showSettings=FALSE, verbose=FALSE)
```

### Arguments

<code>stringMin</code>	vector with minimum values for each gene.
<code>stringMax</code>	vector with maximum values for each gene.
<code>suggestions</code>	optional list of suggested chromosomes
<code>popSize</code>	the population size.
<code>iters</code>	the number of iterations.
<code>mutationChance</code>	the chance that a gene in the chromosome mutates. By default $1/(size+1)$ . It affects the convergence rate and the probing of search space: a low chance results in quicker convergence, while a high chance increases the span of the search space.
<code>elitism</code>	the number of chromosomes that are kept into the next generation. By default is about 20% of the population size.
<code>monitorFunc</code>	Method run after each generation to allow monitoring of the optimization
<code>evalFunc</code>	User supplied method to calculate the evaluation function for the given chromosome
<code>showSettings</code>	if true the settings will be printed to screen. By default False.
<code>verbose</code>	if true the algorithm will be more verbose. By default False.

## References

C.B. Lucasius and G. Kateman (1993). Understanding and using genetic algorithms - Part 1. Concepts, properties and context. *Chemometrics and Intelligent Laboratory Systems* 19:1-33.

C.B. Lucasius and G. Kateman (1994). Understanding and using genetic algorithms - Part 2. Representation, configuration and hybridization. *Chemometrics and Intelligent Laboratory Systems* 25:99-145.

## See Also

[rbga.bin plot.rbga](#)

## Examples

```
# optimize two values to match pi and sqrt(50)
evaluate <- function(string=c()) {
  returnVal = NA;
  if (length(string) == 2) {
    returnVal = abs(string[1]-pi) + abs(string[2]-sqrt(50));
  } else {
    stop("Expecting a chromosome of length 2!");
  }
  returnVal
}

monitor <- function(obj) {
  # plot the population
  xlim = c(obj$stringMin[1], obj$stringMax[1]);
  ylim = c(obj$stringMin[2], obj$stringMax[2]);
  plot(obj$population, xlim=xlim, ylim=ylim,
       xlab="pi", ylab="sqrt(50)");
}

rbga.results = rbga(c(1, 1), c(5, 10), monitorFunc=monitor,
                  evalFunc=evaluate, verbose=TRUE, mutationChance=0.01)

plot(rbga.results)
plot(rbga.results, type="hist")
plot(rbga.results, type="vars")
```

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rbga.bin

*R Based Genetic Algorithm (binary chromosome)*

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## Description

A R based genetic algorithm that optimizes, using a user set evaluation function, a binary chromosome which can be used for variable selection. The optimum is the chromosome for which the evaluation value is minimal.

It requires a `evalFunc` method to be supplied that takes as argument the binary chromosome, a vector of zeros and ones. Additionally, the GA optimization can be monitored by setting a `monitorFunc` that takes a `rbga` object as argument.

Results can be visualized with `plot.rbga` and summarized with `summary.rbga`.

## Usage

```
rbga.bin(size=10,  
         suggestions=NULL,  
         popSize=200, iters=100,  
         mutationChance=NA,  
         elitism=NA, zeroToOneRatio=10,  
         monitorFunc=NULL, evalFunc=NULL,  
         showSettings=FALSE, verbose=FALSE)
```

## Arguments

<code>size</code>	the number of genes in the chromosome.
<code>popSize</code>	the population size.
<code>iters</code>	the number of iterations.
<code>mutationChance</code>	the chance that a gene in the chromosome mutates. By default $1/(size+1)$ . It affects the convergence rate and the probing of search space: a low chance results in quicker convergence, while a high chance increases the span of the search space.
<code>elitism</code>	the number of chromosomes that are kept into the next generation. By default is about 20% of the population size.
<code>zeroToOneRatio</code>	the change for a zero for mutations and initialization. This option is used to control the number of set genes in the chromosome. For example, when doing variable selectionm this parameter should be set high to
<code>monitorFunc</code>	Method run after each generation to allow monitoring of the optimization
<code>evalFunc</code>	User supplied method to calculate the evaluation function for the given chromosome
<code>showSettings</code>	if true the settings will be printed to screen. By default False.
<code>verbose</code>	if true the algorithm will be more verbose. By default False.
<code>suggestions</code>	optional list of suggested chromosomes

## References

C.B. Lucasius and G. Kateman (1993). Understanding and using genetic algorithms - Part 1. Concepts, properties and context. *Chemometrics and Intelligent Laboratory Systems* 19:1-33.

C.B. Lucasius and G. Kateman (1994). Understanding and using genetic algorithms - Part 2. Representation, configuration and hybridization. *Chemometrics and Intelligent Laboratory Systems* 25:99-145.

## See Also

[rbga plot.rbga](#)

**Examples**

```

# a very simplistic optimization
evaluate <- function(string=c()) {
  returnVal = 1 / sum(string);
  returnVal
}

rbga.results = rbga.bin(size=10, mutationChance=0.01, zeroToOneRatio=0.5,
  evalFunc=evaluate)

plot(rbga.results)

# in this example the four variables in the IRIS data
# set are complemented with 36 random variables.
# Variable selection should find the four original
# variables back (example by Ron Wehrens).
## Not run:
data(iris)
library(MASS)
X <- cbind(scale(iris[,1:4]), matrix(rnorm(36*150), 150, 36))
Y <- iris[,5]

iris.evaluate <- function(indices) {
  result = 1
  if (sum(indices) > 2) {
    huhn <- lda(X[,indices==1], Y, CV=TRUE)$posterior
    result = sum(Y != dimnames(huhn)[[2]][apply(huhn, 1,
      function(x)
        which(x == max(x)))]]) / length(Y)
  }
  result
}

monitor <- function(obj) {
  minEval = min(obj$evaluations);
  plot(obj, type="hist");
}

woppa <- rbga.bin(size=40, mutationChance=0.05, zeroToOneRatio=10,
  evalFunc=iris.evaluate, verbose=TRUE, monitorFunc=monitor)

## End(Not run)

# another realistic example: wavelength selection for PLS on NIR data
## Not run:
library(pls.pcr)
data(NIR)

numberOfWavelengths = ncol(NIR$Xtrain)
evaluateNIR <- function(chromosome=c()) {
  returnVal = 100
  minLV = 2

```

```

    if (sum(chromosome) < minLV) {
      returnVal
    } else {
      xtrain = NIR$Xtrain[,chromosome == 1];
      pls.model = pls(xtrain, NIR$Ytrain, validation="CV", grpsize=1,
                     ncomp=2:min(10,sum(chromosome)))
      returnVal = pls.model$val$RMS[pls.model$val$nLV-(minLV-1)]
      returnVal
    }
  }
}

monitor <- function(obj) {
  minEval = min(obj$evaluations);
  filter = obj$evaluations == minEval;
  bestObjectCount = sum(rep(1, obj$popSize)[filter]);
  # ok, deal with the situation that more than one object is best
  if (bestObjectCount > 1) {
    bestSolution = obj$population[filter,][1,];
  } else {
    bestSolution = obj$population[filter,];
  }
  outputBest = paste(obj$iter, " #selected=", sum(bestSolution),
                    " Best (Error=", minEval, "): ", sep="");
  for (var in 1:length(bestSolution)) {
    outputBest = paste(outputBest,
                      bestSolution[var], " ",
                      sep="");
  }
  outputBest = paste(outputBest, "\n", sep="");

  cat(outputBest);
}

nir.results = rbga.bin(size=numberOfWavelengths, zeroToOneRatio=10,
                      evalFunc=evaluateNIR, monitorFunc=monitor,
                      popSize=200, iters=100, verbose=TRUE)

## End(Not run)

```

---

summary.rbga

*R Based Genetic Algorithm Summary Function*


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## Description

Summarizes the genetic algorithm results.

## Usage

```
## S3 method for class 'rbga'
summary(object, echo=FALSE, ...)
```

**Arguments**

object	a rbga object.
echo	if true, the summary will be printed to STDOUT as well as returned.
...	other options (ignored)

**Examples**

```
evaluate <- function(string=c()) {  
  returnVal = 1 / sum(string);  
  returnVal  
}
```

```
rbga.results = rbga.bin(size=10, mutationChance=0.01, zeroToOneRatio=0.5,  
  evalFunc=evaluate)
```

```
summary(rbga.results)
```



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