

Package ‘gramEvol’

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Title Grammatical Evolution for R

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Description A native R implementation of grammatical evolution (GE).
GE facilitates the discovery of programs that can achieve a desired goal.
This is done by performing an evolutionary optimisation over a population
of R expressions generated via a user-defined context-free grammar (CFG)
and cost function.

URL <https://github.com/fnoorian/gramEvol/>

BugReports <https://github.com/fnoorian/gramEvol/issues>

Depends

Suggests rex, knitr

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c	<i>Grammar Rule Concatenation</i>
---	-----------------------------------

Description

Concatenates two or more grammar rule objects.

Usage

```
## S3 method for class 'GERule'
c(..., recursive=FALSE)
```

Arguments

...	Grammar rule objects to be concatenated.
recursive	Not used.

Value

A new grammar rule object.

See Also

[CreateGrammar](#)

Examples

```
rule1 <- grule(Func1, Func2)
rule2 <- grule(`*`, `\/`)

rule.all <- c(rule1, rule2)

print(rule.all)
```

CreateGrammar *Context-free Grammar Object*

Description

Creates a context-free grammar object.

Usage

```
grule(...)
```

```
gsrule(...)
```

```
gvrule(vec)
```

```
CreateGrammar(ruleDef, startSymb)
```

Arguments

...	A series of comma separated strings or expressions, for <code>gsrule</code> and <code>grule</code> respectively. Expressions can be wrapped in <code>.()</code> to preserve their commas or assignment operators.
<code>vec</code>	An iterable vector or list.
<code>ruleDef</code>	Grammatical rule definition. Either a list of grammar rule objects (<code>GERule</code>) created using <code>grule</code> and <code>gsrule</code> with a syntax similar to Backus-Naur form, or a list of character strings representing symbols and sequences in Backus-Naur form, or a filename or connection to a <code>.bnf</code> file. See details.
<code>startSymb</code>	The symbol where the generation of a new expression should start. If not given, the first rule in <code>ruleDef</code> is used.

Details

The rule definition is the grammar described in Backus-Naur context-free grammatical format. The preferred way of defining a grammar is to create a list simulating BNF format, which collects several named grammar rule objects (`GERule`). Each name defines the *non-terminal symbol*, and each rule in the collection determines the *production rule*, i.e., possible *sequences* that will replace the symbol.

Defining a grammar rule object (`GERule`) can take three forms:

1. The first form uses `grule` (Grammar Rule), where R expressions are accepted. In the mapping process, variables are looked up and replaced using the production rules.
2. The second form uses `gsrule` (Grammar String Rule) and uses character strings. The input to `gsrule` are character string values, where any value surrounded by `'<'` or `'>'` is considered as *non-terminal symbols* and will be replaced using the rule with the same name in the mapping process.

Other symbols are considered terminals. This form allows generation of sequences that are not syntactically valid in R (such as ``var op var``).

3. The third form uses `gvrule` (Grammar Vector Rule), where objects within an iterable (vector or list) containing all of the expressions are used as individual rules.

Alternatively, `CreateGrammar` can read and parse `.bnf` text files.

Value

`CreateGrammar` returns a grammar object.

`grule` and `gsrule` return a `GERule` object.

See Also

[c](#), [GrammarMap](#), [GrammaticalEvolution](#)

Examples

```
# Define a simple grammar in BNF format
# <expr> ::= <var><op><var>
# <op>   ::= + | - | *
# <var>  ::= A | B
ruleDef <- list(expr = gsrule("<var><op><var>"),
               op   = gsrule("+", "-", "*"),
               var  = gsrule("A", "B"))

# print rules
print(ruleDef)

# create and display a vector rule
vectorRule = gvrule(1:5)
print(vectorRule)

# Create a grammar object
grammarDef <- CreateGrammar(ruleDef)

# print grammar object
print(grammarDef)

# Creating the same grammar using R expressions
ruleDef <- list(expr = grule(op(var, var)),
               op   = grule(`+`, `-`, `*`),
               var  = grule(A, B))

grammarDef <- CreateGrammar(ruleDef)

print(grammarDef)

# Two rules with commas and assignments, preserved using .()
ruleDef <- list(expr = grule(data.frame(dat)),
               dat  = grule.(x = 1, y = 2), .(x = 5, y = 6)))
```

```
grammarDef <- CreateGrammar(ruleDef)
print(GrammarMap(c(0), grammarDef))
print(GrammarMap(c(1), grammarDef))
```

EvalExpressions	<i>Evaluate a collection of Expressions</i>
-----------------	---

Description

EvalExpressions evaluates one or more expressions, either in string format or as expression objects.

Usage

```
EvalExpressions(expressions, envir = parent.frame())
```

Arguments

expressions	an expression, or a collection of expressions.
envir	the environment in which expressions are to be evaluated. May also be NULL, a list, a data frame, a pair-list or an integer as specified in <code>sys.call</code> .

Details

EvalExpressions is a wrapper around `eval` and `parse` functions in R base package. It can handle a single, a vector or a list of expressions, character strings or GEPHENOTYPE objects.

The `envir` argument is directly passed to `eval` function. If it is not specified, the parent frame (i.e., the environment where the call to `eval` was made) is used instead.

EvalExpressions only evaluates terminal expressions and character strings. Evaluating non-terminal expressions will result in a warning and NA is returned.

Value

If one expression is evaluated, a vector of numeric values is returned. Otherwise a data frame with result of each expression in a separate column is returned.

See Also

[GrammarMap](#)

Examples

```

A <- 1:6
B <- 1

EvalExpressions("A - B")

# a vector of text strings
exprs <- c("A + B", "A - B")
EvalExpressions(exprs, data.frame(A = A, B = B))

# a vector of expressions
exprs <- expression(A + B, A - B)
EvalExpressions(exprs, data.frame(A = A, B = B))

```

EvolutionStrategy.int *Evolution Strategy with Integer Chromosomes*

Description

Uses evolution strategy to find the minima of a given cost function. It evolves chromosomes with limited-range integers as codons.

Usage

```

EvolutionStrategy.int(genomeLen, codonMin, codonMax,
  genomeMin = rep.int(codonMin, genomeLen),
  genomeMax = rep.int(codonMax, genomeLen),
  suggestion = NULL, popSize=4, newPerGen = 4,
  iterations = 500, terminationCost = NA,
  mutationChance = 1/(genomeLen+1),
  monitorFunc = NULL, evalFunc, allowrepeat = TRUE,
  showSettings = FALSE, verbose = FALSE, plapply = lapply)

```

Arguments

genomeLen	Number of integers (i.e, codons) in chromosome.
codonMin	Minimum integer value range for all codons.
codonMax	Maximum integer value range for all codons.
genomeMin	A vector of length genomeLen containing fine-grained control over each codon's minimum. Overrides codonMin.
genomeMax	A vector of length genomeLen containing fine-grained control over each codon's maximum. Overrides codonMax.
suggestion	A list of suggested chromosomes to be used in the initial population.
popSize	Size of the population generated by mutating the parent.
newPerGen	Number of the new randomly generated chromosome in each generation.

<code>iterations</code>	Number of generations to evolve the population.
<code>terminationCost</code>	Target cost. If the best chromosome's cost reaches this value, the algorithm terminates.
<code>mutationChance</code>	The chance of a codon being mutated. It must be between 0 and 1.
<code>monitorFunc</code>	A function that is called at each generation. Can be used to monitor evolution of population.
<code>evalFunc</code>	The cost function.
<code>allowrepeat</code>	Allows or forbids repeated integers in the chromosome.
<code>showSettings</code>	Enables printing GA settings.
<code>verbose</code>	Enables verbose debugging info.
<code>plapply</code>	<code>lapply</code> function used for mapping chromosomes to the cost function. See details for parallelization tips.

Details

`EvolutionStrategy.int` implements evolutionary strategy search algorithm with chromosomes created from integer values in the range of `codonMin` to `codonMax`. `genomeMin` and `genomeMax` allow fine-grained control of range for individual codons. It first creates an initial population, using suggested input suggestion or a randomly generated chromosome. Score of each chromosome is evaluated using the cost function `costFunc`. If the best chromosome reaches `terminationCost`, the algorithm terminates; otherwise only the best candidate is selected and mutated to create a new generation, and the cycle is repeated. This iteration continues until the required cost is reached or the number of generations exceeds `iterations`.

At each generation, the supplied `monitorFunc` is called with a list similar to `EvolutionStrategy.int` returning value as its argument.

The `evalFunc` receives integer sequences and must return a numeric value. The goal of optimization would be to find a chromosome which minimizes this value.

To parallelize cost function evaluation, set `plapply` to a parallelized `lapply`, such as `mclapply` from package `parallel`. In functions that do not handle data dependencies such as `parLapply`, variables and functions required for correct execution of `evalFunc` must be exported to worker nodes before invoking `EvolutionStrategy.int`.

Value

A list containing information about settings, population, and the best chromosome.

<code>settings\$genomeMin</code>	Minimum of each codon.
<code>Settings\$genomeMax</code>	Maximum of each codon.
<code>settings\$popSize</code>	Size of the population created using mutation.
<code>settings\$newPerGen</code>	Number of the new randomly generated chromosome in each generation.

```

settings$totalPopulation      Size of the total population.
settings$iterations          Number of maximum generations.
settings$suggestion          Suggested chromosomes.
settings$mutationChance      Mutation chance.
population$population        The genomic data of the current population.
population$evaluations       Cost of the latest generation.
population$best              Historical cost of the best chromosomes.
population$mean              Historical mean cost of population.
population$currentIteration  Number of generations evolved until now.
best$genome                  The best chromosome in integer sequence format.
best$cost                    The cost of the best chromosome.

```

See Also

[GrammaticalEvolution](#), [GeneticAlg.int](#)

Examples

```

# define the evaluate function
evalfunc <- function(l) {
  # maximize the odd indices and minimize the even indices
  # no repeated values are allowed
  odd <- seq(1, 20, 2)
  even <- seq(2, 20, 2)
  err <- sum(l[even]) - sum(l[odd]);

  stopifnot(!any(duplicated(l))) # no duplication allowed

  return (err)
}

monitorFunc <- function(result) {
  cat("Best of gen: ", min(result$best$cost), "\n")
}

x <- EvolutionStrategy.int(genomeLen = 20, codonMin = 0, codonMax = 20,
  allowrepeat = FALSE, terminationCost = -110,
  monitorFunc = monitorFunc, evalFunc = evalfunc)

```



```

print(x)

best.result <- x$best$genome
print("Odds:")
print(sort(best.result[seq(1, 20, 2)]))
print("Evens:")
print(sort(best.result[seq(2, 20, 2)]))

```

GeneticAlg.int

Genetic Algorithm with Integer Chromosomes

Description

Uses genetic algorithm to find the minima of a given cost function. It evolves chromosomes with limited-range integers as codons.

Usage

```

GeneticAlg.int(genomeLen, codonMin, codonMax,
  genomeMin = rep.int(codonMin, genomeLen),
  genomeMax = rep.int(codonMax, genomeLen),
  suggestions = NULL, popSize = 50,
  iterations = 100, terminationCost = NA,
  mutationChance = 1/(genomeLen+1), elitism = floor(popSize/10),
  geneCrossoverPoints = NULL,
  monitorFunc = NULL, evalFunc, allowrepeat = TRUE,
  showSettings = FALSE, verbose = FALSE, plapply = lapply)

```

Arguments

genomeLen	Number of integers (i.e, codons) in chromosome.
codonMin	Minimum integer value range for all codons.
codonMax	Maximum integer value range for all codons.
genomeMin	A vector of length genomeLen containing fine-grained control over each codon's minimum. Overrides codonMin.
genomeMax	A vector of length genomeLen containing fine-grained control over each codon's maximum. Overrides codonMax.
suggestions	A list of suggested chromosomes to be used in the initial population. Alternatively, an m-by-n matrix, where m is the number of suggestions and n is the chromosome length.
popSize	Size of the population.
iterations	Number of generations to evolve the population.
terminationCost	Target cost. If the best chromosome's cost reaches this value the algorithm terminates.

mutationChance	The chance of a codon being mutated. It must be between 0 and 1.
geneCrossoverPoints	Codon groupings (genes) to be considered while crossover occurs. If given, odd and even codon groups are exchanged between parents. Otherwise random points are selected and a classic single-point crossover is performed.
elitism	Number of top ranking chromosomes that are directly transferred to next generation without going through evolutionary operations.
monitorFunc	A function that is called at each generation. Can be used to monitor evolution of population.
evalFunc	The cost function.
allowrepeat	Allows or forbids repeated integers in the chromosome.
showSettings	Enables printing GA settings.
verbose	Enables verbose debugging info.
plapply	lapply function used for mapping chromosomes to cost function. See details for parallelization tips.

Details

GeneticAlg.int implements evolutionary algorithms with chromosomes created from integer values in the range of codonMin to codonMax. genomeMin and genomeMax allow fine-grained control of range for individual codons. It first creates an initial population, using suggested inputs suggestions and randomly generated chromosomes. Cost of each chromosome is evaluated using the cost function evalFunc. If one of the chromosomes reaches terminationCost, the algorithm terminates; Otherwise evolutionary operators including selection, cross-over and mutation are applied to the population to create a new generation. This iteration is continued until the required cost is reached or the number of generations exceeds iterations.

At each generation, the supplied monitorFunc is called with a list similar to GeneticAlg.int returning value as its argument.

The evalFunc receives integer sequences and must return a numeric value. The goal of optimization would be to find a chromosome which minimizes this value.

To parallelize cost function evaluation, set plapply to a parallelized lapply, such as mclapply from package parallel. In functions that do not handle data dependencies such as parLapply, variables and functions required for correct execution of evalFunc must be exported to worker nodes before invoking GeneticAlg.int.

Value

A list containing information about settings, population, and the best chromosome.

settings\$genomeMin	Minimum of each codon.
Settings\$genomeMax	Maximum of each codon.
settings\$popSize	Size of the population.

```

settings$elitism      Number of elite individuals.
settings$iterations  Number of maximum generations.
settings$suggestions Suggested chromosomes.
settings$mutationChance Mutation chance.
settings$geneCrossoverPoints Cross-over points.
population$population The genomic data of the current population.
population$evaluations Cost of the latest generation.
population$best      Historical cost of the best chromosomes.
population$mean      Historical mean cost of population.
population$currentIteration Number of generations evolved until now.
best$genome          The best chromosome.
best$cost            The cost of the best chromosome.

```

References

This function is partially inspired by `genalg` package by Egon Willighagen. See <https://cran.r-project.org/package=genalg>.

See Also

[GrammaticalEvolution](#), [EvolutionStrategy.int](#)

Examples

```

# define the evaluate function
evalfunc <- function(l) {
  # maximize the odd indices and minimize the even indices
  # no repeated values are allowed
  odd <- seq(1, 20, 2)
  even <- seq(2, 20, 2)
  err <- sum(l[even]) - sum(l[odd]);

  stopifnot(!any(duplicated(l))) # no duplication allowed

  return (err)
}

monitorFunc <- function(result) {

```

```

    cat("Best of gen: ", min(result$best$cost), "\n")
  }

x <- GeneticAlg.int(genomeLen = 20, codonMin = 0, codonMax = 20,
  allowrepeat = FALSE, terminationCost = -110,
  monitorFunc = monitorFunc, evalFunc = evalfunc)

print(x)

best.result <- x$best$genome
print("Odds:")
print(sort(best.result[seq(1, 20, 2)]))
print("Evens:")
print(sort(best.result[seq(2, 20, 2)]))

```

GrammarGetNextSequence

Grammar Iterator

Description

Iterates through grammar's valid sequences.

Usage

```

GrammarGetFirstSequence(grammar,
  seqStart = NULL,
  startSymb = GrammarStartSymbol(grammar),
  max.depth = GrammarGetDepth(grammar),
  max.len = GrammarMaxSequenceLen(grammar, max.depth, startSymb))

```

```

GrammarGetNextSequence(grammar,
  seqStart = NULL,
  startSymb = GrammarStartSymbol(grammar),
  max.depth = GrammarGetDepth(grammar),
  max.len = GrammarMaxSequenceLen(grammar, max.depth, startSymb))

```

```
is.GrammarOverflow(object)
```

Arguments

grammar	A grammar object.
seqStart	The sequence to be incremented. For a value of NULL, the first sequence is returned. Partial sequences are completed and returned.
startSymb	The non-terminal symbol where the generation of a new expression should start.
max.depth	Maximum depth of recursion, in case of a cyclic grammar. By default it is limited to the number of production rules in the grammar.

<code>max.len</code>	Maximum length of sequence to return. Used to avoid recursion.
<code>object</code>	An object to be tested.

Details

`GrammarGetFirstSequence` returns the first sequence that creates a valid expression with the given `grammar` object. `GrammarGetNextSequence` allows iterating through all valid sequences in a grammar. If a `seqStart = NULL` is used, `GrammarGetFirstSequence` is called to and the first sequence in the grammar is returned. Calling `GrammarGetNextSequence` or `GrammarGetFirstSequence` with an incomplete sequence returns a full-length sequence starting with the given `seqStart`.

When `GrammarGetNextSequence` reaches the last of all valid sequences, it returns a `GrammarOverflow` object. This object can be identified using `is.GrammarOverflow`.

Value

`GrammarGetFirstSequence` returns a numeric vector representing the first sequence of the grammar.

`GrammarGetNextSequence` returns a numeric vector or a `GrammarOverflow` object.

`is.GrammarOverflow` returns TRUE if `object` is a `GrammarOverflow`, otherwise FALSE.

See Also

[GrammaticalExhaustiveSearch](#)

Examples

```
# Define a simple grammar
# <expr> ::= <var><op><var>
# <op>   ::= + | - | *
# <var>  ::= A | B
ruleDef <- list(expr = gsrule("<var><op><var>"),
               op   = gsrule("+", "-", "*"),
               var  = gsrule("A", "B"))

# Create a grammar object
grammarDef <- CreateGrammar(ruleDef)

# Iterate and print all valid sequence and expressions
string <- NULL
while (TRUE) {
  string <- GrammarGetNextSequence(grammarDef, string)

  if (is.GrammarOverflow(string)) {
    break
  }

  expr <- GrammarMap(string, grammarDef)
  cat(string, " -> ", as.character(expr), "\n")
}
```

```
# test a partial string
GrammarGetNextSequence(grammarDef, c(0, 0, 2))
```

GrammarIsTerminal *Non-terminal Phenotype test.*

Description

Checks a phenotype object for containing non-terminal symbols.

Usage

```
GrammarIsTerminal(x)
```

Arguments

x A [GEPhenotype](#) object.

Value

TRUE if phenotype is terminal, FALSE otherwise.

See Also

[GrammarMap](#)

Examples

```
# Define a recursive grammar
# <expr> ::= <expr>+<expr> | var
# <var> ::= A | B | C
ruleDef <- list(expr = grule(expr+expr, var),
               var = grule(A, B, C))

# Create a grammar object
grammarDef <- CreateGrammar(ruleDef)

# a short sequence leading to infinite recursion
sq <- c(0)
expr <- GrammarMap(sq, grammarDef)

print(expr)

# check the phenotype for being non-terminal
print(GrammarIsTerminal(expr))

# a terminal sequence
sq <- c(0, 1, 0, 1, 2)
```

```

expr <- GrammarMap(sq, grammarDef)

print(expr)
print(GrammarIsTerminal(expr))

```

GrammarMap

*Sequence to Expression Mapping using Context-free Grammar***Description**

Converts a sequence of integer numbers to an expression using a grammar object.

Usage

```
GrammarMap(inputString, grammar, wrappings = 3, verbose = FALSE)
```

Arguments

inputString	A vector of integers to define the path of symbol selection in grammar tree. It uses zero-based indexing to address production rules in the grammar.
grammar	A grammar object.
wrappings	The number of times the function is allowed to wrap around inputString if non-terminal symbols are still remaining.
verbose	Prints out each steps of grammar mapping.

Details

GrammarMap starts from the startExpr defined in the [grammar](#) object; then it iterates through inputString, replacing symbols in the expression with associated replacements in the grammar using the current value of inputString.

If the function exhausts all non-terminal symbols in the expression, it terminates. If the end of inputString is reached and still non-terminal symbols exist, the algorithm will restart from the beginning of the current inputString. To avoid unlimited recursions in case of a cyclic grammar, wrappings variable limits the number of this restart.

If verbose = TRUE, step-by-step replacement of symbols with production rules are displayed.

GrammarMap returns a GEPhenotype object, which can be converted to a character string using as.character, or an R expression with as.expression.

Value

A GrammarMap returns a GEPhenotype object.

expr	The generated expression as a character string.
parsed	The generated expression. NULL if the expression still contains non-terminal symbols.
type	"T" if the expression is valid, "NT" if the expression still contains non-terminal symbols.


```
wrappings = 3,
retries = 100)
```

Arguments

grammar	A grammar object.
numExpr	Number of random expressions to generate.
max.depth	Maximum depth of recursion, in case of a cyclic grammar. By default it is limited to the number of production rules in the grammar.
startSymb	The symbol where the generation of a new expression should start.
max.string	Maximum value for each element of the sequence.
wrappings	The number of times the function is allowed to wrap around <code>inputString</code> if non-terminal symbols are still remaining.
retries	Number of retries until a terminal and valid expressions is found.

Details

GrammarRandomExpression creates `num.expr` random expressions from the given grammar. It can be used to quickly examine the expressibility of the grammar, or as a form of random search over the grammar.

Value

An expressions, or a list of expressions.

Examples

```
# Define a simple grammar
# <expr> ::= <var><op><var>
# <op>   ::= + | - | *
# <var>  ::= A | B | C
ruleDef <- list(expr = gsrule("<var><op><var>"),
               op =  gsrule("+", "-", "*"),
               var =  grule(A, B, C))

# Create a grammar object
grammarDef <- CreateGrammar(ruleDef)

# Generate 5 random expressions
exprs <- GrammarRandomExpression(grammarDef, 5)
print(exprs)
```

 GrammaticalEvolution *Grammatical Evolution*

Description

Evolves an expression using a context-free grammar to minimize a given cost function.

Usage

```
GrammaticalEvolution(grammarDef, evalFunc,
  numExpr = 1,
  max.depth = GrammarGetDepth(grammarDef),
  startSymb = GrammarStartSymbol(grammarDef),
  seqLen = GrammarMaxSequenceLen(grammarDef, max.depth, startSymb),
  wrappings = 3,
  suggestions = NULL,
  optimizer = c("auto", "es", "ga"),
  popSize = "auto", newPerGen = "auto", elitism = 2,
  mutationChance = NA,
  iterations = "auto",
  terminationCost = NA,
  monitorFunc = NULL,
  disable.warnings=FALSE,
  plapply = lapply, ...)

## S3 method for class 'GrammaticalEvolution'
print(x, ..., show.genome = FALSE)
```

Arguments

<code>grammarDef</code>	A grammar object.
<code>evalFunc</code>	The cost function, taking a string or a collection of strings containing the expression(s) as its input and returning the cost of the expression(s).
<code>numExpr</code>	Number of expressions generated and given to <code>evalFunc</code> .
<code>max.depth</code>	Maximum depth of search in case of a cyclic grammar. By default it is limited to the number of production rules in the grammar.
<code>startSymb</code>	The symbol where the generation of a new expression should start.
<code>seqLen</code>	Length of integer vector used to create the expression.
<code>wrappings</code>	Number of wrappings in case the length of chromosome is not enough for conversion to an expression.
<code>suggestions</code>	Suggested chromosomes to be added to the initial population pool. if <code>optimizer</code> parameter is set to "es", only a single chromosome (as a numeric vector) is acceptable. For "ga" mode, a list of numeric vectors.

optimizer	The evolutionary optimizer. "es" uses evolution strategy as in EvolutionStrategy.int and "ga" uses genetic algorithm as in GeneticAlg.int . "auto" chooses evolution strategy when numExpr = 1, and genetic algorithm otherwise. If "auto" is used, popSize and iterations are tweaked based on the grammar as well.
popSize	Population size in the evolutionary optimizer. By default, 8 for ES and 48 for GA.
newPerGen	Number of randomly generated individuals in evolution strategy. If "auto", it is set to 25% of population of grammar if it is not recursive, otherwise to all of it.
elitism	Number of top ranking chromosomes that are directly transferred to the next generation without going through evolutionary operations, used in genetic algorithm optimizer.
iterations	Number of maximum iterations in the evolutionary optimizer. By default, 1000 for "es" optimizer and 200 for "ga".
terminationCost	Target cost. If a sequence with this cost or less is found, the algorithm terminates.
mutationChance	Mutation chance in the evolutionary optimizer. It must be between 0 and 1. By default it is set to $1/(1+\text{chromosome size})$ for genetic algorithm and $10/(1+\text{chromosome size})$ for evolution strategy.
monitorFunc	A function that is called at each generation. It can be used to monitor evolution of population.
disable.warnings	If TRUE, suppresses any warnings generated while evaluating evalFuncs.
lapply	lapply function used for mapping chromosomes to the cost function. See details for parallelization tips.
...	Additional parameters are passed to GeneticAlg.int or EvolutionStrategy.int .
x	Grammatical Evolution results.
show.genome	Prints the numeric value of genome if TRUE.

Details

This function performs an evolutionary search over the grammar, better known as Grammatical Evolution. It evolves integer sequences and converts them to a collection containing numExpr expression. These expressions can be evaluated using eval function. The evalFunc receives these expressions and must return a numeric value. The goal of optimization would be to find a chromosome which minimizes this function.

Two evolutionary optimizers are supported: Genetic algorithm and evolution strategy, which are set by the optimizer parameter.

Only valid (i.e., terminal) expressions are passed to evalFunc, and it is guaranteed that evalFunc receives at least one expression.

If the grammar contains recursive elements, it is advisable that chromosomeLen is defined manually, as in such cases the possible search space grows explosively with the recursion. The evolutionary algorithm automatically removes the recursive chromosomes from the population by imposing a penalty for chromosomes creating expressions with non-terminal elements.

monitorFunc receives a list similar to the GrammaticalEvolution's return value.

Value

The results of [GeneticAlg.int](#) or [EvolutionStrategy.int](#) with an additional item:

```
best$expressions
      Expression(s) with the best cost.
```

See Also

[CreateGrammar](#), [GeneticAlg.int](#), [EvolutionStrategy.int](#), [EvalExpressions](#)

Examples

```
# Grammar Definition
ruleDef <- list(expr      = gsrule("<der.expr><op><der.expr>"),
               der.expr = grule(func(var), var),
               func     = grule(log, exp, sin, cos),
               op       = gsrule("+", "-", "*"),
               var      = grule(A, B, n),
               n        = grule(1, 2, 3, 4))

# Creating the grammar object
grammarDef <- CreateGrammar(ruleDef)

# cost function
evalFunc <- function(expr) {
  # expr: a string containing a symbolic expression
  # returns: Symbolic regression Error
  A <- 1:6
  B <- c(2, 5, 8, 3, 4, 1)

  result <- eval(as.expression(expr))

  X <- log(A) * B
  err <- sum((result - X)^2)

  return(err)
}

# invoke grammatical evolution (with default parameters)
ge <- GrammaticalEvolution(grammarDef, evalFunc, terminationCost = 0.001)

# print results
print(ge, sequence = TRUE)
```

GrammaticalExhaustiveSearch

Exhaustive Search

Description

Exhaustive Search within context-free grammar.

Usage

```
GrammaticalExhaustiveSearch(grammar, evalFunc,
    max.depth = GrammarGetDepth(grammar),
    startSymb = GrammarStartSymbol(grammar),
    max.len = GrammarMaxSequenceLen(grammar, max.depth, startSymb),
    wrappings = 3,
    terminationCost = NA,
    monitorFunc = NULL)
```

Arguments

grammar	A grammar object.
evalFunc	The evaluation function, taking an expression as its input and returning the cost (i.e., the score) of the expression.
max.depth	Maximum depth of recursion, in case of a cyclic grammar. By default it is limited to the number of production rules in the grammar.
startSymb	The symbol where the generation of a new expression should start.
max.len	Maximum length of the sequences to search. By default it is determined by max.depth.
wrappings	The number of times the function is allowed to wrap around inputString if non-terminal symbols are still remaining.
terminationCost	Target cost. If an expression with this cost or less is found, the algorithm terminates.
monitorFunc	A function that is called at each iteration. It can be used to monitor the search.

Details

GrammaticalExhaustiveSearch performs an exhaustive search, iterating through all possible expressions that can be generated by the grammar, to find the expression that minimises evalFunc.

The search terminates when all possible expressions are exhausted, or when an expression with a cost less than terminationCost is discovered.

If a monitorFunc is given, it is called for each expression, and it receives a list similar to the GrammaticalExhaustiveSearch's return value with the information available for that expression.

Value

bestExpression	The Best expression.
bestSequence	Best expression's generating sequence.
bestCost	Best expression's cost.
numExpr	Number of evaluated expressions.

In addition, the monitorFunc receives the following additional slots:

currentExpression	The current expression.
-------------------	-------------------------

currentSequence Current expression's generating sequence.
 currentCost Current expression's cost.

See Also

[GrammarGetNextSequence](#), [GrammaticalEvolution](#)

Examples

```
library("gramEvol")

ruleDef <- list(expr = gsrule("<var><op><var>"),
               op  = gsrule("+", "-", "*"),
               var  = gsrule("A", "B"))

# Create a grammar object
grammarDef <- CreateGrammar(ruleDef)

# use exhaustive search to find the sequence for creating "B - A"
evalFunc <- function(expr) {
  if (as.character(expr) == "B - A") {
    return(0) # Minimum error
  } else {
    return(1) # maximum error
  }
}

res <- GrammaticalExhaustiveSearch(grammarDef, evalFunc, terminationCost = 0)

print(res)
```

GrammaticalRandomSearch

Random Search

Description

Random Search within context-free grammar.

Usage

```
GrammaticalRandomSearch(grammar, evalFunc,
                        max.depth = GrammarGetDepth(grammar),
                        startSymb = GrammarStartSymbol(grammar),
                        wrappings = 3,
```

```
iterations = 1000,
terminationCost = NA,
monitorFunc = NULL)
```

Arguments

grammar	A grammar object.
evalFunc	The evaluation function, taking an expression as its input and returning the cost (i.e., the score) of the expression.
max.depth	Maximum depth of recursion, in case of a cyclic grammar. By default it is limited to the number of production rules in the grammar.
startSymb	The symbol where the generation of a new expression should start.
wrappings	The number of times the function is allowed to wrap around inputString if non-terminal symbols are still remaining.
iterations	Number of random expressions to test.
terminationCost	Target cost. If an expression with this cost or less is found, the algorithm terminates.
monitorFunc	A function that is called at each generation. It can be used to monitor evolution of population.

Details

GrammaticalRandomSearch performs a random search within expressions that can be generated by the grammar, to find the expression that minimises evalFunc.

The search terminates when either the predetermined number of iterations are reached, or when an expression with a cost less than terminationCost is discovered.

If a monitorFunc is given, it is called for each expression, and it receives a list similar to the GrammaticalExhaustiveSearch's return value with the information available for that expression.

Value

bestExpression	The Best expression.
bestSequence	Best expression's generating sequence.
bestCost	Best expression's cost.
numExpr	Number of evaluated expressions.
population	A matrix of sequences that were tested.
populationCost	Numeric value of cost of sequences that were tested.

In addition, the monitorFunc receives the following additional slots:

currentExpression	The current expression.
currentSequence	Current expression's generating sequence.
currentCost	Current expression's cost.

See Also

[GrammarGetNextSequence](#), [GrammaticalEvolution](#)

Examples

```
library("gramEvol")

ruleDef <- list(expr = gsrule("<var><op><var>"),
               op   = gsrule("+", "-", "*"),
               var  = gsrule("A", "B"))

# Create a grammar object
grammarDef <- CreateGrammar(ruleDef)

# use exhaustive search to find the sequence for creating "B - A"
evalFunc <- function(expr) {
  if (as.character(expr) == "B - A") {
    return(0) # Minimum error
  } else {
    return(1) # maximum error
  }
}

# search and terminate after getting to cost = 0
res <- GrammaticalRandomSearch(grammarDef, evalFunc, terminationCost = 0)

print(res)
```

ReplaceInExpression *Replace as sub-expression inside an expression*

Description

Replace every subexpression equal to or starting with what in `expr`. Replacement is performed by passing the whole subexpression to `replacer.func`, which should be a function(`x`, ...), where `x` is the expression and return the desirable expression.

Usage

```
ReplaceInExpression(expr, what, replacer.func, ...)
```

Arguments

<code>expr</code>	An expression .
<code>what</code>	A backquoted expression to find in <code>expr</code> .
<code>replacer.func</code>	A function(<code>x</code> , ...) to process the subexpression.
...	Other parameters passed to <code>replacer.func</code> .

Details

This function was designed to be used as a runtime processing tool for grammar generated expression. This allows the user to modify the resulting expression on the fly based on runtime variables, without including them in the grammar. See examples section.

Value

An expression

References

See <http://adv-r.had.co.nz/Expressions.html> by Hadley Wickham.

Examples

```
expr = expression(function(x) {
  cbind(f1(x),
        f2(x),
        g3(y))
})
expr
ReplaceInExpression(expr, bquote(f2), function(x) {NULL})
ReplaceInExpression(expr, bquote(f2), function(x) {bquote(f2(y))})
ReplaceInExpression(expr, bquote(g3), function(x) {bquote(f3(x))})
ReplaceInExpression(expr, bquote(g3), function(x, b) {if (b > 1) x else NULL}, b = 0)
ReplaceInExpression(expr, bquote(g3), function(x, b) {if (b > 1) x else NULL}, b = 2)
```

summary

Context-free Grammar Object Information

Description

Examines a context-free grammar object.

Usage

```
## S3 method for class 'grammar'
summary(object, ...)

GrammarStartSymbol(grammar)

GrammarIsRecursive(grammar, startSymb = GrammarStartSymbol(grammar), ...)

GrammarGetDepth(grammar, max.depth = max(length(grammar$def), 4),
  startSymb = GrammarStartSymbol(grammar), ...)

GrammarMaxSequenceLen(grammar, max.depth = GetGrammarDepth(grammar),
```

```
startSymb = GrammarStartSymbol(grammar), ...)
```

```
GrammarMaxRuleSize(grammar)
```

```
GrammarMaxSequenceRange(grammar, max.depth = GrammarGetDepth(grammar),
startSymb = GrammarStartSymbol(grammar), approximate = FALSE, ...)
```

```
GrammarNumOfExpressions(grammar, max.depth = GrammarGetDepth(grammar),
startSymb = GrammarStartSymbol(grammar), ...)
```

Arguments

grammar, object	A grammar object.
max.depth	Maximum depth of search in case of a cyclic grammar. By default it is limited to the maximum of 4 or the number of production rules in the grammar.
startSymb	The symbol where the generation of a new expression should start.
approximate	If True, results are approximated. Useful for recursive grammars, where number of valid expressions prohibits an accurate measurement.
...	unused inputs.

Value

summary returns a `summary.grammar` object, with the following slots which are obtained from the other functions:

Start.Symbol	GrammarStartSymbol returns the grammar's starting symbol.
Is.Recursive	GrammarIsRecursive returns TRUE if grammar contains a recursive element.
Tree.Depth	GrammarGetDepth returns the depth of the grammar. It is limited to max.depth for a recursive grammar.
Maximum.Sequence.Length	GrammarMaxSequenceLen returns the maximum length of a sequence needed to generate an expression without wrapping.
Maximum.Rule.Size	GrammarMaxRuleSize returns the largest rule size in the grammar.
Maximum.Sequence.Variation	GrammarMaxSequenceRange returns a numeric sequence, with each of its elements holding the highest range that the same position in all sequences can hold.
No.of.Unique.Expressions	GrammarNumOfExpressions returns the number of expressions a grammar can generate.

See Also

[CreateGrammar](#), [GrammarMap](#)

Examples

```
# Define a simple grammar
# <expr> ::= <var><op><var>
# <op>   ::= + | - | *
# <var>  ::= A | B
ruleDef <- list(expr = gsrule("<var><op><var>"),
               op   = gsrule("+", "-", "*"),
               var  = gsrule("A", "B"))

# Create a grammar object
grammarDef <- CreateGrammar(ruleDef)

# summarize grammar object
summary(grammarDef)
```

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