

Package ‘RFreak’

November 8, 2009

Version 0.2-6

Date 2009-09-07

Title R/FrEAK interface

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Depends R (>= 2.6.0), rJava (>= 0.5.0)

Imports methods

Description An R interface to a modified version of the Free Evolutionary Algorithm Kit FrEAK. FrEAK is a toolkit written in Java to design and analyze evolutionary algorithms. Both the R interface and an extended version of FrEAK are contained in the RFreak package. For more information on FrEAK see <http://sourceforge.net/projects/freak427/>.

LazyLoad yes

License GPL (>= 2)

SystemRequirements Java (>= 5.0)

Repository CRAN

Date/Publication 2009-11-08 10:24:18

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<code>data.logicfs</code>	<i>Example Data of GPAS</i>
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Description

`data.logicfs` contains two objects: a simulated matrix `data.logicfs` of 400 observations (rows) and 15 variables (columns) and a vector `cl.logicfs` of length 400 containing the class labels of the observations.

Each variable is categorical with realizations 1, 2 and 3. The first 200 observations are cases, the remaining are controls. If one of the following expressions is TRUE, then the corresponding observation is a case:

`SNP1 == 3`

`SNP2 == 1 AND SNP4 == 3`

`SNP3 == 3 AND SNP5 == 3 AND SNP6 == 1`

where `SNP1` is in the first column of `data.logicfs`, `SNP2` in the second, and so on.

<code>evolreg-class</code>	<i>Class "evolreg"</i>
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Description

Encapsulates information returned from FrEAK computing robust regression. For compatibility reasons also called `ltsEA`.

Objects from the Class

An `evolreg` object holds four slots.

Slots

summary: A `data.frame` with a summary of the FrEAK run (inherited from "`FreakReturn`")

best: The best subset found

coefficients: Vector of coefficient estimates

crit: The value of the objective function of the used regression method

Extends

Class "`FreakReturn`", directly.

Author(s)

Robin Nunkesser <Robin.Nunkesser@tu-dortmund.de>

See Also

["FreakReturn"](#), [robreg.evol](#)

executeSchedule	<i>Executes a FrEAK schedule</i>
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Description

Executes a schedule created by the FrEAK schedule editor and returns a summary of the result.

Usage

```
executeSchedule(freakfile = "schedule.freak")
```

Arguments

`freakfile` File containing the schedule to be executed

Value

Returns an object of class `FreakReturn` enwrapping a `data.frame` in its only slot `summary` containing information about the last population of the executed schedule. For each individual in the last population the following information is contained:

<code>run</code>	The run the individual was found in
<code>generation</code>	The generation the individual was created in
<code>objective value(s)</code>	The objective value(s) as returned by the fitness function
<code>individual</code>	A string representation of the individual

Warning

To obtain a result, the schedule needs to have a stopping criterion and the observer "Result" and the view "R Return" which are automatically preselected when using `launchScheduleEditor`.

Author(s)

Robin Nunkesser <Robin.Nunkesser@tu-dortmund.de>

See Also

[launchScheduleEditor](#), ["FreakReturn"](#)

Examples

```
## Not run:
# Start the schedule editor and set up a schedule
launchScheduleEditor()

# Execute the set up schedule.
executeSchedule()

## End(Not run)
```

FreakReturn-class *Class "FreakReturn"*

Description

Encapsulates information returned from FrEAK

Objects from the Class

A `FreakReturn` object holds only one slot, containing a `data.frame` with the summary of an executed FrEAK run.

Slots

`summary`: A `data.frame` with informations for each returned individual on the run the individual was found in, the generation the individual was created in, the objective value(s) as returned by the fitness function, and the individual itself.

Author(s)

Robin Nunkesser <Robin.Nunkesser@tu-dortmund.de>

See Also

"ltsEA", "GPAS"

GPAS-class *Class "GPAS"*

Description

Encapsulates information returned from FrEAK computing the GPAS algorithm.

Objects from the Class

A `GPAS` object holds two slots.

Slots

`summary`: A `data.frame` with a summary of the FrEAK run (inherited from "[FreakReturn](#)")
`trees`: The Java objects representing the individuals returned by the FrEAK run

Methods

predict Method to obtain predictions of an individual based on new predictors

Extends

Class "[FreakReturn](#)", directly.

Author(s)

Robin Nunkesser <Robin.Nunkesser@tu-dortmund.de>

See Also

["FreakReturn"](#), [GPASDiscrimination](#), [GPASInteractions](#)

GPASDiscrimination *Execute the GPAS algorithm for discrimination*

Description

Working on categorical data with binary response, the algorithm searches for multi-valued logic expressions in disjunctive normal form discriminating between response 0 and response 1. The algorithm is intended for genetic association studies on SNP data.

Usage

```
GPASDiscrimination(resp.train, preds.train, resp.test=NULL,  
  preds.test=NULL, runs = 1, generations = 10000)
```

Arguments

<code>resp.train</code>	Vector with the response variables of the training data set
<code>preds.train</code>	Matrix or data frame with the predictors of the training data set
<code>resp.test</code>	Optional vector with the response variables of the test data set
<code>preds.test</code>	Optional matrix or data frame with the predictors of the test data set
<code>runs</code>	Number of independent runs of GPAS
<code>generations</code>	Number of generations after which the algorithm will be stopped

Value

Returns an object of class GPAS with a `data.frame` in its slot `summary` containing information about the last population of the executed discrimination runs. For each individual in the last population the following information is contained:

<code>data set</code>	Either 'training' or 'test' or omitted
<code>run</code>	The run the individual was found in
<code>generation</code>	The generation the individual was created in
<code>objective value 1</code>	Correctly predicted cases
<code>objective value 2</code>	Correctly predicted controls
<code>objective value 3</code>	Length of the individual
<code>individual</code>	A string representation of the individual

Author(s)

Robin Nunkesser <Robin.Nunkesser@tu-dortmund.de>

References

R. Nunkesser, T. Bernholt, H. Schwender, K. Ickstadt, and I. Wegener (2007). Detecting High-Order Interactions of Single Nucleotide Polymorphisms Using Genetic Programming. *Bioinformatics*, **23**, 3280-3288.

See Also

"GPAS", [GPASInteractions](#)

Examples

```
# load example data
data(data.logicfs)

# execute GPAS to discriminate between cases and controls
GPASDiscrimination(cl.logicfs,data.logicfs,runs=1,generations=1000)
```

GPASInteractions *Execute the GPAS algorithm for feature selection*

Description

Identification of interesting (high order) SNP interactions. The algorithm works on categorical data with binary response and delivers multi-valued logic expressions in disjunctive normal form typically explaining subsets of the data and an interaction tree containing interesting interactions.

Usage

```
GPASInteractions(resp, preds, runs = 1, generations = 10000,  
  savegraph = "interactions.dot", occurences=10, ratio=0.1)
```

Arguments

<code>resp</code>	Vector with the response variables
<code>preds</code>	Matrix or data frame with the predictors
<code>runs</code>	Number of independent runs of GPAS
<code>generations</code>	Number of generations after which the algorithm will be stopped
<code>savegraph</code>	Name of the file the resulting GraphViz graph will be saved to
<code>occurences</code>	The minimum number of times an interaction has to occur to be included in the graph
<code>ratio</code>	The minimal ratio a single literal has to occur in relation to his ancestor in the interaction graph to be included in the interaction graph

Value

Returns an object of class `GPAS` with a `data.frame` in its slot `summary` containing information about the last population of the executed runs. For each individual in the last population the following information is contained:

<code>run</code>	The run the individual was found in
<code>generation</code>	The generation the individual was created in
<code>objective value 1</code>	Sum of correctly predicted cases and controls
<code>objective value 2</code>	Correctly predicted controls
<code>objective value 3</code>	Length of the individual
<code>individual</code>	A string representation of the individual

Author(s)

Robin Nunkesser <Robin.Nunkesser@tu-dortmund.de>

References

R. Nunkesser, T. Bernholt, H. Schwender, K. Ickstadt, and I. Wegener (2007). Detecting High-Order Interactions of Single Nucleotide Polymorphisms Using Genetic Programming. *Bioinformatics*, **23**, 3280-3288.

GraphViz: <http://www.graphviz.org/>

See Also

"GPAS", [GPASDiscrimination](#)

Examples

```
# load example data
data(data.logicfs)

# execute GPAS to search for interesting interactions
GPASInteractions(cl.logicfs,data.logicfs,runs=1,generations=1000)
```

```
launchScheduleEditor
```

Launches a graphical schedule editor

Description

Launches a graphical schedule editor for FrEAK schedules. A schedule contains the algorithm and simulation options for the desired evolutionary algorithm.

Usage

```
launchScheduleEditor(saveTo = "schedule.freak", load = NULL)
```

Arguments

saveTo	Name of the file the schedule is saved to
load	Name of an (optional) file that should be loaded to the editor

Details

Setting up a schedule consists of seven algorithm specific steps and two simulation specific steps. To set up an evolutionary algorithm it is necessary to choose a search space, a fitness function, an optional genotype-mapper, an algorithm graph, a set of stopping criteria, a population model, and an initial population. The simulation specific steps allow the user to choose views and observers (the necessary ones for the R interface are preselected, views depending on the FrEAK GUI are not supported). The last step consists of choosing batches (only one batch supported) and the number of independent runs. A detailed instruction on how to set up schedules can be found in the User's Guide of FrEAK.

Warning

Editing schedules has to be finished by pressing the "Finish" button to obtain a file including the schedule. The observer "Result" and the view "R Return" - which are preselected - and a stopping criterion are necessary for the R interface to work.

Author(s)

Robin Nunkesser <Robin.Nunkesser@tu-dortmund.de>

References

FrEAK User's Guide:

<http://ls2-www.cs.uni-dortmund.de/~nunkesser/software/usersguide.pdf>

See Also

[executeSchedule](#)

Examples

```
## Not run:
# Start the schedule editor and set up a schedule
launchScheduleEditor()

## End(Not run)
```

LTSevol

Least Trimmed Squares Robust Regression

Description

Carries out least trimmed squares (LTS) robust regression with an evolutionary algorithm. The LTS regression method minimizes the sum of the h smallest squared residuals. Deprecated. Use [robreg.evol](#) instead.

Usage

```
## Deprecated:
LTSevol(y, x, h = NULL, adjust = FALSE, runs = 1, generations = 10000)
```

Arguments

<code>y</code>	Vector with the response variables
<code>x</code>	Matrix or data frame containing the explanatory variables
<code>h</code>	Parameter determining the trimming
<code>adjust</code>	Whether to perform intercept adjustment at each step
<code>runs</code>	Number of independent runs
<code>generations</code>	Number of generations after which the algorithm will be stopped

Value

The function `LTSevol` returns an object of class "ItsEA". This object contains:

<code>summary</code>	Summary of the FrEAK run
<code>best</code>	The best subset found
<code>coefficients</code>	Vector of coefficient estimates
<code>crit</code>	The value of the objective function of the LTS regression method, i.e., the sum of the h smallest squared raw residuals

Author(s)

Robin Nunkesser <Robin.Nunkesser@tu-dortmund.de>

References

O. Morell, T. Bernholt, R. Fried, J. Kunert, and R. Nunkesser (2008). An Evolutionary Algorithm for LTS-Regression: A Comparative Study. *Proceedings of COMPSTAT 2008*. To Appear.

P. J. Rousseeuw (1984), Least Median of Squares Regression. *Journal of the American Statistical Association* **79**, 871–881.

See Also

"ltsEA"

Examples

```
# load example data
data(stackloss)

# compute LTS regression
LTSevol(stackloss[, 4], stackloss[, 1:3], adjust=TRUE, runs=1, generations=1000)
```

predict-method *Predict using new predictors*

Description

Takes an individual from a GPAS object and predicts on the basis of new predictors.

Usage

```
## S4 method for signature 'GPAS':
predict(object, individual, preds)
```

Arguments

object	Object of class GPAS
individual	Number of the individual to use
preds	New predictors

Value

Returns a vector with the new predictions.

Author(s)

Robin Nunkesser <Robin.Nunkesser@tu-dortmund.de>

See Also

"GPAS", [GPASDiscrimination](#), [GPASInteractions](#)

robreg.evol

Robust Evolutionary Regression

Description

Carries out robust regression with an evolutionary algorithm. `ltsreg`, `ltareg`, `lmsreg`, `lqsreg`, and `lqdreg` are wrappers.

Usage

```
robreg.evol(x, y, method = c("lts", "lta", "lms", "lqs", "lqd"),
           quantile=NULL, adjust=FALSE, runs=1, generations=10000, duration=0)
```

```
## Wrappers:
ltsreg.evol(...)
ltareg.evol(...)
lmsreg.evol(...)
lqsreg.evol(...)
lqdreg.evol(...)
```

Arguments

<code>x</code>	Matrix or data frame containing the explanatory variables
<code>y</code>	Vector with the response variables
<code>method</code>	The method to be used. One of "lts", "lta", "lms", "lqs", and "lqd".
<code>quantile</code>	The quantile to be used: see Details .
<code>adjust</code>	Whether to perform intercept adjustment at each step
<code>runs</code>	Number of independent runs
<code>generations</code>	Number of generations after which the algorithm will be stopped
<code>duration</code>	Duration in seconds after which the algorithm will be stopped
<code>...</code>	Arguments to be passed to the default method

Details

Suppose there are n data points and p regressors, including any intercept.

The first four methods minimize some function of the sorted squared residuals. For methods "lqs" and "lms" it is the `quantile` squared residual, and for "lts" ("lts") it is the sum of the `quantile` smallest squared (absolute) residuals. "lqd" minimizes approximately the quartile of the absolute residual differences.

Value

The function `robreg.evol` returns an object of class "evolreg". This object contains:

<code>summary</code>	Summary of the FrEAK run
<code>best</code>	The best subset found
<code>coefficients</code>	Vector of coefficient estimates
<code>crit</code>	The value of the objective function of the regression method

Author(s)

Robin Nunkesser <Robin.Nunkesser@tu-dortmund.de>

References

O. Morell, T. Bernholt, R. Fried, J. Kunert, and R. Nunkesser (2008). An Evolutionary Algorithm for LTS-Regression: A Comparative Study. *Proceedings of COMPSTAT 2008*. To Appear.

P. J. Rousseeuw (1984), Least Median of Squares Regression. *Journal of the American Statistical Association* **79**, 871–881.

See Also

["evolreg"](#)

Examples

```
# load example data
data(stackloss)

# compute different regressions
robreg.evol(stackloss[, 1:3], stackloss[, 4], method= "lts", generations=1000)
lqsreg.evol(stackloss[, 1:3], stackloss[, 4], generations=1000)
lqdreg.evol(stackloss[, 1:3], stackloss[, 4], generations=1000)
```

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