STAT 243: Model Selection with Genetic Algorithms using GA

Eddie Buehler, Yang Hu & Jin Rou New University of California, Berkeley

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1 Introduction

A genetic algorithm is an optimization technique that mimics the biological process of natural selection. The basic strategy is to take a population of candidate solutions and "evolve" them over many generations towards an optimal solution. Continuing the biology analogy, the population is typically represented as a collection of a series of bits called chromosomes. A fitness function is evaluated at each iteration to determine how well the current population of chromosomes solves the objective function. A genetic algorithm has the following steps:

- 1. Calculate fitness of chromosomes. Often, the fitness is the value of the objective function solving the optimization. The better the fitness, the better solution the chromosomes will provide.
- 2. Select chromosomes to form a mating pool based on their fitness. This process is performed stochastically, with more fit chromosomes having a higher probability of being selected.
- 3. Recombine parent chromosomes from the mating pool. This is the first step in producing the next generation of chromosomes in which the parents' genes are combined.
- 4. Apply mutation to produce the resulting generation of chromosomes. After crossover, each gene in the resulting population has a finite probability to flip its value.
- 5. Repeat until either a set number of generations is reached.

2 Code

We took the S3 approach to object-oriented programming for our package and created functions that were as modular as possible to facilitate code creation, maintenance, and testing.

For a given data set, e.g. the built-in airquality data set in R, the user can carry out model selection for an ordinary linear regression of the variable Ozone on other variables in the data set with the main function in the package as follows:

```
ga <- select(data = airquality, yvar = "Ozone")</pre>
```

Finer control of other parameters in the genetic algorithm for model selection is possible by changing the other function arguments. More details can be found in the GA manual in the Appendix.

The output of the results can be viewed using the following commands:

```
summary(ga)
model(ga)
```

The result of this function is an object of GA class that contains the settings, model data, final population of chromosomes/models, model evaluation values of this population and all results of model selection using the genetic algorithm.

In the genetic algorithmic approach to model selection, a population of models, i.e. chromosomes with number of genes equal to the number of model variables under consideration, and with each gene taking a value of 1 if the model variable is included and 0 otherwise, is first initialized. This population then undergoes many iterations of reproduction. In the reproduction stage, each model/chromosome is then evaluated based on the desired model selection criterion, e.g. Akaike's Information Criterion. Chromosomes are then selected into the mating pool based on how well they perform on the model selection criterion.

Next, 2 parent chromosomes are randomly selected from the mating pool to form a child chromosome, with some probability of recombination/crossover occurring in the process. This is repeated until the desired number of child chromosomes for the next generation is reached. Finally, mutation is applied with a low probability to each gene in the population of child chromosomes. The resulting population of child chromosomes forms the next generation. The whole process is repeated for a desired number of iterations. The model/chromosome with the minimum value of the model evaluation criterion across all generations is then the best model.

The following subsections elaborate on the main subfunctions in the GA package; for more detailed documentation, please refer to the GA manual in the Appendix.

initialize function

We sample uniformly from the set of $\{0, 1\}$ with replacement as many genes as is required, i.e. the product of the number of variables under consideration and the population size desired, for the initial population of chromosomes.

evaluate function

We compute the value of the model evaluation criterion for every model/chromosome in the population. The default model evaluation criterion is Akaike Information Criterion (AIC). The user can also choose to use Bayesian Information Criterion (BIC) or define his/her own function.

select_for_mating function

The default selection method is "rank", which refers to Linear Rank Selection (LRS). In LRS, chromosomes are first given ranks r_i for i = 1, ..., n, where n is the number of chromosomes in the population. The chromosome with the best (minimum) model evaluation criterion is assigned a rank of n. Chromosomes are then selected into the mating pool randomly with probability proportional to their relative rank, until the desired size of the mating pool is reached. This is done with the following algorithm:

- 1. Calculate for each chromosome its probability to be selected, $p_i = \frac{r_i}{\sum\limits_{i=1}^n r_i}$. Since the best chromosome is given the largest rank, it also has a highest probability of being selected.
- 2. Calculate the cumulative probability for each chromosome to be selected, $pc_i = \sum_{j=1}^{i} p_j$.
- 3. Generate a random number u uniformly in the range [0, 1].
- 4. Select the chromosome with index i if $pc_i < u < pc_{i+1}$
- 5. Repeat until the desired number of chromosomes in the mating pool is selected.

The alternative selection method is "tournament", which refers to Tournament Selection. The idea is simple; in Tournament Selection, we randomly select 2 chromosomes from the population. The chromosome with the better model evaluation criterion is selected into the mating pool. This process is repeated (with replacement of both chromosomes each time) until the desired number of chromosomes in the mating pool is reached.

recombine function

Recombination occurs with probability prob_recombine for every set of 2 parent chromosomes; the child chromosome is simply a copy of the first parent chromosome if no recombination occurs. We implemented three methods of recombination: crossover at one point ("onepoint"), crossover at two points ("twopoint"), and uniform crossover ("uniform").

For one-point crossover, a break point index b is uniformly sampled from the set of $\{1, ..., g\}$, where g is the number of genes on each chromosome. The resulting child chromosome then takes genes 1 to b from the first parent and genes b+1 to g from the second parent.

For two-point crossover, two break point indices b_1 and b_2 are uniformly sampled without replacement from the set of $\{1, ..., g\}$. The resulting child chromosome then takes genes 1 to b_1 from the first parent, genes $b_1 + 1$ to b_2 from the second parent, and finally genes $b_2 + 1$ to g from the first parent.

For uniform crossover, the child chromosome has equal probability of receiving each gene from either parent.

mutate function

We generate as many Bernoulli variables as there are genes in the population of chromosomes. For each gene, we set:

$$\text{new_gene} \sim \begin{cases} Bernoulli(\texttt{prob_mutate}), & \text{if current_gene} = 0 \\ Bernoulli(1-\texttt{prob_mutate}), & \text{if current_gene} = 1 \end{cases}$$

This is equivalent to mutating each gene (i.e. changing a value of 1 to 0 and vice versa) with a probability of prob_mutate, but doing it in this manner allows for vectorized operations.

3 Testing

Automated testing of auxilliary functions was performed using R's testthat package. This approach has several advantages over manual testing. It provides functions that allow the user to clearly define expected inputs, outputs, and errors while testing. Additionally, re-testing simply involves running the test file again, and multiple test files can be run at once. Finally, it displays test progress visually, showing whether each tests passes, fails, or produces an error.

Each function was tested for correct inputs and outputs. The number of inputs, their class, and their

dimensions was specified using the expect_that(x, equals(y)) and expect_that(x, is_a(y)) functions. Similarly, the number of outputs, their class, and their dimensions were specified using the same functions. Further constraints on certain inputs and outputs were tested as well. Since chromosomes are composed of bits, the elements in the population matrix must be either zero or one. This specification was tested by ensuring that the minimum of the population is 0 and the maximum is 1.

Although most functions had stochastically determined outputs and thus their exact values could not be predicted, the mutate function had two testable outcomes. First, if the mutation rate was set to zero, no genes were expected to mutate, so the output population should be the exact same as the input population. Secondly, if the mutation rate were one, all of the genes in the population should switch their value. Both of these scenarios were tested to ensure proper behavior in the mutate function.

A testing function was written for each function in the ga package. These files can be found in the /testing folder of the Github repository. All testing files can be run at once using the test_dir(''~/testing'') command in R.

After ensuring that each function takes proper inputs and returns desired outputs, each method was also tested using small data sets to see if they functions properly. In order to test the functionality of our genetic algorithm, we employed a larger, more realistic data set. We compared the models selected using our genetic algorithm with a well-known model selection method, the stepwise model selection using AIC, implemented in R in the stepAIC function available in the MASS package.

This data set was obtained from surveys about how video games affect grades. There are 15 variables in the data set – time (number of hours played), like (whether like to play), where (where to play), freq (how often), busy (play if busy), educ (playing educational), sex, age, home (computer at home), math (hate math), work (number of hours work per week), own (own PC), cdrom (PC has CD-rom), email (have Email) and grade. The dependent variable is grade. Completed data were obtained from 91 students during Fall 1994 at Berkeley. The data source can be found at the Stat Labs website for University of California, Berkeley.

The following results are obtained using our genetic algorithm.

```
res <- summary(ga)
```

```
## Model 1 :
## grade ~ where + freq + busy + sex + home + math
## AIC = 157.6
## -----
## Model 2 :
## grade ~ freq + educ + sex + home + math
## AIC = 158.2
## -----
## Model 3 :
## grade ~ freq + busy + sex + home + math
## AIC = 158.3
## -----
## Model 4 :
## grade ~ where + freq + busy + sex + home + math + own
## AIC = 158.4
## -----
## Model 5 :
## grade ~ where + freq + busy + sex + home + math + email
## AIC = 158.6
```

The following results are obtained using the stepAIC function.

```
library (MASS)
mod <- glm(grade ~ ., data = data)</pre>
res_step <- stepAIC(mod)</pre>
## Start: AIC=167.2
## grade ~ time + like + where + freq + busy + educ + sex + age +
      home + math + work + own + cdrom + email
##
##
            Df Deviance AIC
## - age 1 23.6 165
## - where 1
                    23.6 166
## - time 1
                    23.6 166
## - like 1 23.7 166
## - educ 1 23.8 166

## - cdrom 1 23.8 166

## - work 1 23.9 166

## - email 1 23.9 167

## - busy 1 23.9 167

## <none> 23.6 167
## - math 1 24.2 168
```

```
## - own 1 24.2 168
## - freq 1 24.6 169
## - home 1
               25.8 174
## - sex 1
               27.4 179
##
## Step: AIC=165.3
## grade ~ time + like + where + freq + busy + educ + sex + home +
## math + work + own + cdrom + email
##
##
        Df Deviance AIC
## - where 1
              23.7 164
## - time
         1
              23.7 164
## - like 1
              23.8 164
## - educ 1
              23.8 164
             23.8 164
23.9 165
## - cdrom 1
## - work 1
## - busy 1
              24.0 165
## - email 1
              24.0 165
## <none>
              23.6 165
             24.2 166
24.3 166
## - math 1
## - own 1
## - freq 1
              24.6 167
## - home 1
              25.9 172
## - sex 1
               27.5 178
##
## Step: AIC=163.7
## grade ~ time + like + freq + busy + educ + sex + home + math +
## work + own + cdrom + email
##
##
         Df Deviance AIC
## - like 1
              23.9 162
## - cdrom 1
               23.9 162
## - busy 1
              24.0 163
## - time 1
              24.0 163
## - work 1
              24.0 163
## - educ 1
              24.1 164
## - math 1
              24.2 164
## <none>
              23.7 164
## - email 1
              24.2 164
## - own
        1
              24.4 165
## - freq 1
              24.8 166
            26.1 170
## - home 1
## - sex 1 27.9 177
##
```

```
## Step: AIC=162.4
## grade ~ time + freq + busy + educ + sex + home + math + work +
    own + cdrom + email
##
         Df Deviance AIC
## - cdrom 1 24.1 161
## - time
               24.2 162
         1
## - work 1
              24.2 162
## - busy 1
              24.2 162
## - math 1
              24.3 162
## - email 1
              24.4 162
              24.4 162
## - educ 1
## <none>
               23.9 162
## - own 1
              24.6 163
## - freq 1
              25.0 164
## - home 1
              26.2 169
## - sex 1
              27.9 175
##
## Step: AIC=161.2
## grade ~ time + freq + busy + educ + sex + home + math + work +
     own + email
##
         Df Deviance AIC
##
## - time 1 24.4 160
## - work 1
               24.4 160
## - busy 1
              24.5 161
## - math 1
              24.5 161
## - educ 1
               24.6 161
## <none>
               24.1 161
## - email 1
              24.6 161
## - own 1
              24.7 162
## - freq 1
              25.1 163
              26.6 168
## - home 1
## - sex 1
               28.6 175
## Step: AIC=160.3
## grade ~ freq + busy + educ + sex + home + math + work + own +
##
     email
##
##
         Df Deviance AIC
## - work 1 24.7 160
## - busy 1
               24.8 160
## - email 1
              24.8 160
              24.9 160
## - educ 1
```

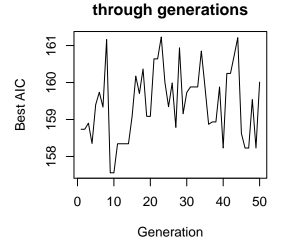
```
## - math 1 24.9 160
## <none>
                24.4 160
## - own
                25.0 161
## - freq 1
                25.6 163
## - home 1
                27.1 168
## - sex
         1
                28.7 173
##
## Step: AIC=159.6
## grade ~ freq + busy + educ + sex + home + math + own + email
##
##
          Df Deviance AIC
                25.1 159
## - email 1
## - busy
                25.2 159
         1
## - own
          1
                25.2 159
## - educ 1
               25.2 159
## <none>
               24.7 160
## - math 1
               25.8 162
## - freq 1
               25.9 162
## - home 1
                27.3 167
## - sex 1
                28.8 171
##
## Step: AIC=159.2
## grade ~ freq + busy + educ + sex + home + math + own
##
        Df Deviance AIC
##
## - own
         1
               25.5 159
## - educ 1
               25.6 159
## - busy 1
               25.6 159
               25.1 159
## <none>
## - math 1
              26.2 161
## - freq 1
              26.3 161
## - home 1
               27.8 166
## - sex 1
               29.2 171
##
## Step: AIC=158.6
## grade ~ freq + busy + educ + sex + home + math
##
        Df Deviance AIC
##
## - busy 1
              26.0 158
## - educ 1
               26.0 158
## <none>
               25.5 159
## - math 1
               26.6 160
## - freq 1
              26.7 161
## - home 1
               27.8 164
```

```
## - sex
           1
                 29.4 169
##
## Step:
         AIC=158.2
## grade ~ freq + educ + sex + home + math
##
##
          Df Deviance AIC
## <none>
                 26.0 158
## - freq 1
                 26.7 159
## - math 1
                 26.8 159
## - educ
                 27.4 161
## - home
                 28.4 164
                 29.8 168
## - sex
```

The best model found using genetic algorithm was: $y \sim \text{where} + \text{freq} + \text{busy} + \text{sex} + \text{home} + \text{math}$, with an AIC of 157.56. This result is better than that of 158.23 that we obtained using the stepAIC function.

Finally, we plotted the best AIC for each generation to see how the best AIC has changed over generations.

```
par(cex = 0.8)
plot(ga)
```



Evolution of best model

From the plot, we can see that the best model was found at generation 9.

4 Contributions

4.1 Code writing

General structure: JRN Functions: JRN, YH

4.2 Code testing

Function testing: EB

Overall function tests: YH

4.3 Documentation

Manual creation: EB

Project write-up: Introduction (EB), Code (JRN), Testing (EB, YH)

5 Appendix

GA-package

Genetic algorithm for model variable selection

Description

Final project for Statistics 243. An R package that implements a genetic algorithm for variable selection in linear and GLM problems.

Details

Package: GA-package

Type: Package

Version: 1.0

Date: 2014-12-13

Author(s)

Eddie Buehler, Yang Hu, JR New

References

G. Givens and J. Hoeting. Computational Statistics, 2nd ed. (2012).

See Also

https://github.com/jrnew/genetic-algo

Examples

```
# Select regression variables for airquality data using lm model and AIC criterion
ga <- select(data = airquality,</pre>
             yvar = "Ozone",
             xvars = NULL,
             model = "lm",
             criterion = "AIC",
             pop_size = 100L,
             method_select = "rank",
             method_recombine = "onepoint",
             prob_recombine = 0.6,
             prob_mutate = 0.01,
             num_max_iterations = 100L,
             seed = 123,
             do_parallel = FALSE)
# With a user-defined model evaluation criterion function
rsquared <- function(lm) {</pre>
  mod <- summary(lm)</pre>
  return(-mod\$r.squared)
ga <- select(data = airquality,</pre>
             yvar = "Ozone",
             model = "lm",
             criterion = "rsquared",
             criterion_function = rsquared)
```

Carry out model selection with a genetic algorithm.

select

Description

Main function for carrying out model selection with a genetic algorithm.

Usage

```
select(data, yvar, xvars = NULL, model = "lm", glm_family = NULL,
    criterion = "AIC", pop_size = 100L, method_select = "rank",
    method_recombine = "onepoint", prob_recombine = 0.6, prob_mutate = 0.01,
    num_max_iterations = 100L, seed = 123, do_parallel = FALSE)
```

Arguments

prob_mutate

data Data frame		
yvar Character; Name of column containing response variable		
xvars Character vector; Default is all column names that are no column(s) containing set of explanatory variables to select	, , ,	
model Character; "lm" (default) or "glm"; Linear model or gener	ralized linear model.	
glm_family Character if model is "glm", NULL otherwise; "binomia fault), "Gamma", "inverse.gaussian", "poisson", "quasi", "son"; A family function that gives the error distribution a be used in the model.	quasibinomial", "quasipois-	
criterion "AIC" (default) or "BIC"; Criterion to be minimized.		
pop_size Integer; Default is 100; Number of chromosomes per gener	ration.	
method_select		
String; "rank" (linear rank selection) (default) or "tourn select chromosomes for inclusion in mating pool.	nament"; Method to	
method_recombine		
String; "onepoint" (default), "twopoint", "uniform"; Type point, at two points or uniformly (at all possible points).	e of crossover, at one	
prob_recombine		
Numeric, between 0 and 1; Default is 0.6; Probability of re	ecombination.	

Numeric, between 0 and 1; Default is 0.01; Probability of mutation.

num_max_iterations

Non-negative integer; Default is 100; Maximum number of iterations before

algorithm is stopped.

seed Non-negative integer; Default is 123; Random seed for reproducibility.

do_parallel Logical; Default is FALSE; Do in parallel?

evaluate_once

Do evaluation once.

Description

Do evaluation for a chromosome by calculating model selection criterion.

Usage

```
evaluate_once(model_data, xvars_select, model = "lm", glm_family = NULL,
    criterion = "AIC")
```

Arguments

model Character; "lm" (default) or "glm"; Linear model or generalized linear model.

glm_family Character if model is "glm", NULL otherwise; "binomial", "gaussian" (de-

fault), "Gamma", "inverse.gaussian", "poisson", "quasi", "quasibinomial", "quasipois-

son"; A family function that gives the error distribution and link function to

be used in the model.

criterion "AIC" (default) or "BIC"; AIC or BIC.

model_data; Object of class model_data.

xvars_select;

Logical vector;

Value

Numeric; Value of criterion.

evaluate

Do evaluation.

Description

Do evaluation for chromosomes in population by calculating model selection criterion.

Usage

```
evaluate(pop, model_data, model = "lm", glm_family = NULL,
    criterion = "AIC", do_parallel = FALSE)
```

Arguments

pop Matrix of population of chromosomes.

model_data Object of class model_data.

model Character; "lm" (default) or "glm"; Linear model or generalized linear model.

glm_family Character if model is "glm", NULL otherwise; "binomial", "gaussian" (de-

fault), "Gamma", "inverse.gaussian", "poisson", "quasi", "quasibinomial", "quasipois-

son"; A family function that gives the error distribution and link function to

be used in the model.

criterion "AIC" (default) or "BIC"; Criterion to be minimized.

do_parallel Logical; Default FALSE; Do in parallel?

Value

Numeric vector; Evaluation values for all chromosomes in the current generation.

 ${\tt initialize} \qquad \qquad {\tt Initialize} \ \ {\tt first} \ generation \ of \ chromosomes.$

Description

Initialize first generation of chromosomes completely randomly.

Usage

```
initialize(pop_size, num_vars)
```

Arguments

pop_size Non-negative integer; Number of chromosomes in population.

num_vars Non-negative integer; Number of variables in model under consideration/ num-

ber of genes in each chromosome.

Value

A matrix of size pop_size x num_vars with 1's and 0's.

mutate

Mutate genes in the population.

Description

Mutate each gene in the population at a pre-defined rate.

Usage

```
mutate(pop, prob_mutate = 0.01)
```

Arguments

pop Matrix; Population of chromosomes.

prob_mutate Numeric, between 0 and 1; Default is 0.01; Probability of mutation.

Value

Matrix of population of chromosomes that have undergone mutation.

plot.ga

Plots results from the genetic algorithm.

Description

Plots the best model evaluation criterion in each generation against the generation iteration.

Usage

```
## S3 method for class 'ga'
plot(ga, num_view = 3)
```

Arguments

ga Object of class ga.

num_view Number of top models to display.

Value

Prints summary of top models and associated value of model selection criterion.

 $process_data$

Process data for input into genetic algorithm.

Description

Process data for input into genetic algorithm.

Usage

```
process_data(data, yvar, xvars = NULL)
```

Arguments

data Data frame

yvar Character; Name of column containing response variable.

xvars Character vector; Default is all column names that are not yvar; Name(s) of

column(s) containing set of explanatory variables to select on.

Value

A list object named model_data containing:

data Data frame; Processed data with only relevant columns.

yvar Character; Name of column containing response variable.

xvars Character vector; Name(s) of column(s) containing set of explanatory variables to select on.

num_vars Integer; Length of xvars.

recombine_once Recombine once.

Description

Carry out crossover of two parent chromosomes to produce one child chromosome.

Usage

recombine_once(parent1, parent2, method = "onepoint")

Arguments

parent1	Integer vector of 1st parent chromosome containing 1's and 0's.
parent2	Integer vector of 2nd parent chromosome containing 1's and 0's.

method String; "onepoint" (default), "twopoint", "uniform"; Type of crossover, at one

point, at two points or uniformly (at all possible points).

Value

Integer vector of child chromosome containing 1's and 0's.

recombine Recombine.

Description

Carry out crossover of parent chromosomes in a mating pool.

Usage

```
recombine(pop_mating, pop_size, method = "onepoint", prob_recombine = 0.6,
   do_parallel = FALSE)
```

Arguments

pop_mating Matrix of population of chromosomes that form the mating pool.

pop_size Integer; Number of chromosomes in a generation.

method String; "onepoint" (default), "twopoint", "uniform"; Type of crossover, at one

point, at two points or uniformly (at all possible points).

prob_recombine

Numeric, between 0 and 1; Default is 0.6; Probability of recombination.

do_parallel Logical; Default FALSE; Do in parallel?

Value

Matrix of population of chromosomes resulting from recombination.

reproduce

 $Wrapper\ function\ for\ reproduction\ stage.$

Description

Wrapper function for reproduction stage.

Usage

```
reproduce(ga, iteration, do_parallel = FALSE)
```

Arguments

ga Object of class ga. iteration Iteration number.

Value

Updated ga list object.

select_for_mating Select chromosomes for recombination.

Description

Select chromosomes for recombination based on fitness.

Usage

```
select_for_mating(pop, evaluation, method = "rank", do_parallel = FALSE)
```

Arguments

pop Matrix; Population of chromosomes.

evaluation Numeric vector; Evaluation values of all chromosomes in population.

method String; "rank" (linear rank selection) (default) or "tournament"; Method to

select chromosomes for inclusion in mating pool.

do_parallel Logical; Default FALSE; Do in parallel?

Value

Matrix of population of chromosomes that form the mating pool.

summary.ga

 $Display\ summary\ of\ results\ from\ the\ genetic\ algorithm.$

Description

Outputs the top models selected from the genetic algorithm.

Usage

```
## S3 method for class 'ga'
summary(ga, num_view = 5)
```

Arguments

ga Object of class ga.

num_view Number of top models to display.

Value

Prints summary of top models and associated value of model selection criterion.