Time Complexity Analysis of the Genetic Algorithm Clustering Method

Z. M. NOPIAH, M. I. KHAIRIR, S. ABDULLAH, M. N. BAHARIN, and A. ARIFIN Department of Mechanical and Materials Engineering Universiti Kebangsaan Malaysia 43600 UKM Bangi, Selangor MALAYSIA zmn@eng.ukm.my

Abstract: - This paper presents the time complexity analysis of the genetic algorithm clustering method. The tested feature in the clustering algorithm is the population limit function. For the purpose of the study, segmental kurtosis analysis was done on several segmented fatigue time series data, which are then represented in two-dimensional heteroscaled datasets. These datasets are then clustered using the genetic algorithm clustering method and at the runtime of the algorithm is measured against the number of iterations. Polynomial fitting is used on the runtime data to determine the time complexity of the algorithm. The results of the analysis will be used to determine the significance of including the population limit function in the algorithm for optimal performance.

Key-Words: - Genetic algorithms, fatigue damage, clustering, time complexity, big-O notation, algorithm efficiency.

1 Introduction

In the field of evolutionary computing, the evolutionary principles of survival of the fittest, natural selection and genetic inheritance are abstracted and modeled into algorithms that search for optimal solutions to a problem. The most popular technique in evolutionary computing research has been the genetic algorithm [1-3]. Genetic algorithms (GA) perform meta-heuristic search in complex, large, and multimodal landscapes, and provide near-optimal solutions for objective or fitness functions of optimization problems[4, 5]. GAs and GAbased techniques have been used in fields such as industrial engineering [1] and in optimizing the performance of neural networks, fuzzy systems, production systems, wireless systems and other program structures [2].

A GA-based clustering method was developed for applications in cluster analysis of heteroscaled datasets. Since this method is to be applied numerous times across multiple datasets in numerous iterations, it is necessary that the method performs efficiently in real time in order to consistently produce efficient results without using up too much computing and time resource. This study aims to analyze the time complexity of the GA clustering algorithm by comparing the real time performance of the method with and without a population limit function. The results will be used to determine whether or not the population limit function must be included in the algorithm for optimal performance.

2 Literature Review

2.1 Genetic Algorithms

Most GA methods have at least the following in common: populations of chromosomes, selection according to fitness, crossover to produce new offspring, and random mutation of new offspring [2]. Solutions in GA are encoded as chromosomes which are strings of numbers or characters that represent the values or parameters of the solution to the problem. The chromosomes are commonly encoded as strings of binary, real-valued, integer, octal, or hexadecimal numbers [1]. Each of these types of numbers has their own advantages and disadvantages when used for certain data types or for searching for solutions to certain problems. In this study, real-valued numbers string was selected as the chromosome encoding for the population of potential solutions.

The set of potential solutions to the problem is represented as a population of chromosomes. Initially, a random population is created, which represents different points in the search space of potential solutions [4, 6, 7]. A fitness function assigns a score (fitness) to each chromosome in the current population, which will determine its survival into the next generation. The fitness of a chromosome depends on how well that chromosome can solve the problem at hand [6].

The selection of chromosomes is done on the current population based on the fitness values – chromosomes with higher fitness are more likely to be selected than those with low fitness values. This is mostly done using probabilistic methods; in evolutionary computing researches, the common methods of selection are the roulette wheel, tournament, and rank selection [1, 2, 4]. Selected chromosomes are then included in the next generation of population.

Next the population undergoes the crossover (also called recombination) genetic operator, which selects chromosomes from the population to produce offsprings. Using random selection or any of the previously mentioned selection methods, two parent chromosomes are chosen for crossover operation. Using single-point, two-point, or *N*-point crossover, parts of the gene string in each parent chromosome are swapped to produce two new offspring, which are included in the next generation of population. The process is repeated a number of times, usually according to some user-specified proportional value of the current population.

Random chromosomes from the surviving population are selected for mutation, where some random part of the chromosome's gene is arbitrarily changed. This genetic operation may or may not yield superior offspring, but it ensures that solutions are not trapped in local extrema. Mutation is performed according to some degree of probability, usually very small, so that the GA does not approximate a random search [1].

The process of selection, crossover, and mutation are then repeated on the surviving population, until some terminating criteria is reached, i.e. a maximum number of generations, a minimum change in population fitness, etc. The resulting final population is then considered to be the set of solutions that best solves the problem at hand. The best individual chromosome (the chromosome with the highest fitness value) in the final population is usually determined to be the optimal solution to the problem.

2.2 Time complexity analysis

Time complexity analysis is a part of computational complexity theory that is used to describe an algorithm's use of computational resources; in this case, the worst case running time expressed as a function of its input using big Omicron (big-*O*) notation [8, 9]. The big-*O* notation is used to express the upper bound of the growth rate of a function and is mostly used to describe asymptotic behavior [8].

The big-*O* notation is described using set notation as follows:

$$O(g(n)) = \{ f \mid \exists c > 0, \exists n_0 > 0, \forall n \ge n_0 : 0 \le f \le cg(n) \}$$
(1)

In other words, $f \in (g(n))$ if and only if there exist positive constants c and n_0 such that for all $n \ge n_0$, the inequality $0 \le f \le cg(n)$ is satisfied. We say that f is big O of g(n), or that g(n) is an asymptotic upper bound for f[9]. In terms of time complexity analysis, we use the term $T(n) \in O(g(n))$ and say that the algorithm has order of g(n) complexity. This means that the time taken to compute a problem of size n is in the set of functions described by O(g(n)).

Time complexity analysis can be used to predict the growth behavior of an algorithm and is useful for analyzing and optimizing the real time efficiency of the algorithm [9].

3 Methodology

In this study, several segmented fatigue time series data (see Table 1) were used to test the GA clustering algorithm for real time efficiency. Segmental kurtosis analysis was done on each segmented fatigue data, and the results are represented in two-dimensional heteroscaled datasets.

Table 1: Description of datasets used in the study

Dataset	Description	
SAETRN	SAE transmission test fatigue data	
DK1	Pavé road loading on lower arm suspension	
DDK2	Highway road loading on lower arm	

The GA clustering algorithm is then used on these datasets to cluster fatigue damage segments based on kurtosis and fatigue damage their values. Simultaneously, the number of population of solutions and the running time are recorded while the algorithm is running. For the purpose of this study, the algorithm is set to run until 1000 generations have been produced, which means that the algorithm has iterated 1000 times. The recorded runtime and population growth are then plotted and polynomial fitting is used to estimate the growth function of the running time.

The processes above are then repeated on the heteroscaled datasets after a population limit function is included in the selection process of the GA clustering algorithm. The theoretical value of the population limit of the GA clustering algorithm is evaluated as

$$P_{max} = C_P P_0 \left(1 + r_c \right) \left(1 + r_m \right)$$
(2)

Equation 2 is derived from how the population size grows with every iteration of the GA clustering algorithm. Initially, the population size is a positive integer P_0 . For a worst case scenario, we assume that the whole population was selected in the selection process. This means that P_0 number of solutions is considered for the crossover operation. A portion of the population is selected for the crossover process, which produces additional solutions to be added to the population. Let r_c be the crossover rate where $0 \le r_c \le 1$; the number of additional solutions would be P_0r_c , making the total number of solutions in the population so far to be $P_0 + P_0r_c$.

Next, a portion of this population is selected for the mutation process. Let r_m be the value of the mutation rate, where $0 \le r_m \le 1$; the number of solutions added to the population is $(P_0 + P_0r_c) r_m$; therefore the total number of solutions in the population after the mutation process is $P_0 + P_0r_c + (P_0 + P_0r_c) r_m$.

The expression above is then simplified using factorization as follows:

$$P_0 + P_0 r_c + (P_0 + P_0 r_c) r_m$$

= $P_0 (1 + r_c + (1 + r_c) r_m)$
= $P_0 (1 + r_c) (1 + r_m)$

In order to enable the user to have some amount of control over the maximum population, the expression above is multiplied with a user-defined coefficient of population C_P , which is a positive real number. This results in Equation 2 as expressed above. For the purpose of this study, the values P_0 , C_P , r_c and r_m are set to be 20, 2, 0.4 and 0.1 respectively.

As with the previous set of data, the population size and running time of the algorithm is recorded simultaneously as the population limited GA clustering algorithm is run on the datasets. The recorded runtime and population growth are then plotted and polynomial fitting is used to estimate the growth function of the running time.

The time complexity of the algorithm is then determined from the fitted growth function. The results are then compared for the GA clustered and the population limited GA clustered datasets to determine the significance of including the population limit function in the GA clustering algorithm.

4 Results and Discussion

For the purpose of understanding the population growth and the time complexity of the GA clustering algorithm, plots of population growth and polynomial fitted runtime are observed and compared.

Figure 1 shows how the population size grows with the number of generations or iterations of the GA clustering algorithm. We can see that it is apparent that for all datasets, although the population sizes increase and decrease erratically with the number of iterations, overall they generally exhibit a positive growth behavior. This means that eventually, after some large enough number of generations, the population size continually increases as the number of generation increases. The increase of the size of the population will in turn increase the problem size for the GA clustering algorithm, which will affect the time complexity of the algorithm. Generally, a larger problem size means larger computing time or resource is needed for the algorithm to complete its task.



Fig. 1: Population size versus number of generations for (a) SAETRN, (b) DK1, and (c) DDK2

Figure 2 shows the actual running time (in seconds) versus the number of iterations and the fitted model used to predict the asymptotic behavior of the runtime for each dataset.



Fig. 2: Polynomial fitting of runtime versus number of iterations for (a) SAETRN, (b) DK1, and (c) DDK2

We can see that the algorithm runs in polynomial time of some degree, and the fitted models estimate the order of polynomial time the algorithm runs in for each dataset. Table 2 shows the fitted models for the runtime for each dataset and the estimated orders of polynomial time. For datasets SAETRN and DK1, the time complexity is $O(n^3)$ and for DDK2 the time complexity is $O(n^5)$. This means that for two datasets, the algorithm runs in cubic time and for the other dataset, the algorithm runs in polynomial time of degree 5. This tells us that since the population size grows unboundedly, the problem size also grows unboundedly and therefore a much larger computing resource is needed for each next iteration of the GA clustering algorithm. Consequently, the algorithm's running time also increases in the order of polynomial degree 3 or 5 depending on the data. These polynomial growth rates are undesirable for optimum algorithm efficiency, since larger problem sizes would require significantly longer runtime periods and much larger computing resources.

Table 2: Fitted models and complexity

Dataset	Fitted model	Complexity
SAETRN	$f(x) = p_1 x^3 + p_2 x^2 + p_3 x + p$	$O(n^3)$
	p_4	
DK1	$f(x) = p_1 x^3 + p_2 x^2 + p_3 x + p$	$O(n^3)$
	p_4	
DDK2	$f(x) = p_1 x^5 + p_2 x^4 + p_3 x^3 + p_3 x^3$	$O(n^5)$
	$p_4x^2 + p_5x + p_6$	

Figure 3 shows the population growth when the GA clustering algorithm is modified to include the population limit function in its selection process. We can see that although the population sizes generally increase with the number of generations, the numbers are capped at a particular value P_{max} which can be obtained using Equation 2.

Figure 4 shows the actual running time in seconds versus the number of iterations and the fitted model used to predict the asymptotic behavior of the runtime for each dataset when the GA clustering algorithm is run with the population limit function included in its selection process. Based on the fitted models, it is clear that for all datasets, the time complexity of the algorithm is O(n), which means that the algorithm runs in linear time. This tells us that the running time increases linearly with the number of iterations, and that each iteration requires some constant time to perform. This case is much more desirable over the case where the algorithm runs in polynomial time of degrees larger than 1. Since the population size is capped at some finite value P_{max} , the problem size ceases to grow unboundedly and consequently the computing resource needed for each iteration of the GA clustering algorithm is eventually capped, making the computing time for each iteration constant.









The results of the time complexity analysis shows that the population limit function in the selection process of the GA clustering algorithm has managed to reduce the time complexity of the algorithm from higher degree polynomial time to linear time. This means that by including the population limit function in the GA clustering algorithm, the running time of the algorithm can be significantly reduced and the user will have some degree of control over the complexity of the algorithm in both computing time and resource.

5 Conclusion

Performing time complexity analysis on the GA clustering algorithm has helped us to determine how the algorithm performs in real time as the problem size increases. It has been found that including the population limit function in the selection process of the GA clustering algorithm will reduce the time complexity of the algorithm to linear time. This significant reduction in time complexity will be very useful in future developments of the GA clustering algorithm, particularly for clustering larger datasets in higher dimensions.

6 Acknowledgements

The authors would like to express their gratitude to Universiti Kebangsaan Malaysia and Ministry of Higher Education, through the fund of UKM-GUP-BTT-07-25-152, for supporting these research activities.

References:

- [1] S.N. Sivanandam, and S.N. Deepa, *Introduction to Genetic Algorithms*, New York: Springer-Verlag Berlin Heidelberg, 2008.
- [2] M. Gen and R. Cheng, *Genetic Algorithms and Engineering Optimization*, New York: Wiley-Interscience, 2000.
- [3] Z. Michalewicz, *Genetic Algorithms* + *Data Structures* = *Evolution Programs*, 3rd rev. ext. ed. New York: Springer-Verlag, 1996.
- [4] M. Mitchell, An Introduction to Genetic Algorithms, 5th print, Cambridge, MA: The MIT Press, 1999.
- [5] U. Maulik and S. Bandyopadhyay, Genetic algorithm-based clustering technique. *Pattern Recognition*, Vol. 33, 2000, pp. 1455-1465.
- [6] N.H. Park, C.W. Ahn and R.S. Ramakrishna, Adaptive clustering technique using genetic algorithms. *IEICE Transactions on Information* and Systems, Vol. E88-D (12), 2005, pp. 2880-2882.
- [7] A. Banerjee and S.J. Louis, A recursive clustering methodology using a genetic algorithm. *IEEE Congress on Evolutionary Computation (CEC)*, 2007, pp. 2165-2172.
- [8] Knuth, D. E., Big Omicron and big Omega and big Theta. SIGACT News Vol. 8, No. 2 (Apr. 1976), 1976, pp. 18-24.

[9] Black, P. E., big-O notation, *Dictionary of Algorithms and Data Structures* [online], U.S. National Institute of Standards and Technology, 2008 (accessed 26 November 2009). Available from: http://www.itl.nist.gov/div897/sqg/dads/HTML/bigOnotation.html