

Efficient Numerical Optimization Algorithm Based on New Real-Coded Genetic Algorithm, AREX + JGG, and Application to the Inverse Problem in Systems Biology

Asako Komori¹, Yukihiro Maki^{2,3}, Masahiko Nakatsui⁴, Isao Ono⁵, Masahiro Okamoto^{1,3*}

¹Department of Bioinformatics, Graduate School of Systems Life Sciences, Kyushu University, Fukuoka, Japan

²Faculty of International Communications, Fukuoka International University, Fukuoka, Japan

³Synthetic Systems Biology Research Center, Kyushu University, Fukuoka, Japan

⁴Department of Systems Bioscience for Drug Discovery, Graduate School of Pharmaceutical Sciences, Kyoto University, Kyoto, Japan

⁵Graduate School of Computational Intelligence and Systems Science, Tokyo Institute of Technology, Tokyo, Japan

Email: *okahon@brs.kyushu-u.ac.jp

Received July 10, 2012; revised August 10, 2012; accepted August 17, 2012

ABSTRACT

In Systems Biology, system identification, which infers regulatory network in genetic system and metabolic pathways using experimentally observed time-course data, is one of the hottest issues. The efficient numerical optimization algorithm to estimate more than 100 real-coded parameters should be developed for this purpose. New real-coded genetic algorithm (RCGA), the combination of AREX (adaptive real-coded ensemble crossover) with JGG (just generation gap), have applied to the inference of genetic interactions involving more than 100 parameters related to the interactions with using experimentally observed time-course data. Compared with conventional RCGA, the combination of UNDX (unimodal normal distribution crossover) with MGG (minimal generation gap), new algorithm has shown the superiority with improving early convergence in the first stage of search and suppressing evolutionary stagnation in the last stage of search.

Keywords: Inverse Problem; S-System Formalism; Gene Regulatory Network; System Identification; Real-Coded Genetic Algorithm

1. Introduction

A system is not just an assembly of more than two components which have unique function and show time-variant behavior. Since there are principles that govern at the system-level, mere collection of database of components does not lead to the understanding of system's functional property. The aim of systems biology is to understand how functional properties and behavior of living organisms are brought about by the interactions of their constituents such as genes, proteins, metabolites and so on. The research strategies of systems biology can be divided into the following four fields: 1) System identification: inference of interaction between system components, 2) System analysis: dynamics of time-variant components, 3) System control: control toward the desirable condition of the system, 4) System design: design the system which realizes a certain dynamic or time-variant behavior, which expands to the research field, "Synthetic Biology". These research developments are

strongly supported by mathematical and computational methodologies, such as inference algorithm, statistical analysis, numerical calculation, nonlinear optimization, computer simulation and so on. Especially at the research field of systems identification, in order to infer the interaction among systems components, we have to develop the powerful inferring engine, in which large numbers of real-coded parameter values related to the interactions can be estimated efficiently using experimentally observed time-course data; algorithm of powerful numerical optimization for inverse problem involving more than 100 numbers of numerical parameters.

Organizationally complex systems such as gene regulatory networks and metabolic pathways are composed of many interacting components. In the many cases, the details of the molecular mechanism that govern interactions among system components are not well known, however, how do we represent mathematical model of such complex processes? most of these processes are nonlinear. Description of these processes requires a representation that is general enough to capture the essence

*Corresponding author.

of the experimentally observed response. One of the best approaches that satisfy this requirement is the “S-system [1]” which is a type of power-law formalism because it is based on a particular type of ordinary differential equation in which the component processes are characterized by power-law functions (see Section 2). The S-system is suitable for conceptual modeling and describing organizationally complex systems including loop or cyclic-structure between system components. However, S-system formalism has a major disadvantage; this formalism includes large number of parameters must be estimated. This number is $2n(n+1)$, where n is the number of system components.

Tominaga *et al.* have developed the numerical optimization method based on the simple genetic algorithm (SGA) in S-system formalism [2]. However, the SGA has two problems that early convergence in the fast stage of search and evolutionary stagnation in the last stage of search. Then, real-coded genetic algorithms (RCGAs) [3] have attracted attention as alternative numerical optimization methods to the SGA. One of the crossover operators for RCGA, known as the unimodal normal distribution crossover (UNDX) [4,5], has shown good performance in optimizing various functions including multimodal ones and benchmark functions with epistasis among the parameters. Sato *et al.* have proposed a new generation-alternation model called the minimal generation gap (MGG) [6,7], to improve early convergence in the first stage of search and to suppress evolutionary stagnation in the last stage of search. So far, we have developed efficient numerical optimization method using S-system modeling and RCGAs, with a combination of the UNDX and the MGG. This method can be applied to not only the estimation of gene networks but also the estimation of metabolic regulatory network. Shikata *et al.* have applied this method to the estimation of the amino acid network [8]. However, this method has not been able to estimate 20 amino acids’ network ($n = 20$), because this method converges slowly in the early stage of search, followed by the stagnation of optimization.

Therefore, in order to achieve the estimation of the larger network involving more than 100 real-coded parameters, we have developed a new estimation method using RCGA. The research group of Kobayashi has proposed just generation gap (JGG) [9,10], modified MGG. Akimoto *et al.* have reported that a combination of the adaptive real-coded ensemble crossover (AREX) with JGG showed the superiority in most of the benchmark functions [11]. In this study, we have applied new RCGA, AREX + JGG, to the inverse problem in Systems Biology, that is inference of regulatory network in genetic system and metabolic pathways for realizing experimentally observed time-course data of system components, and have compared with conventional RCGA, UNDX + MGG.

2. Method

2.1. S-System Formalism

The S-system [1] is a type of power-law formalism because it is based on a particular type of ordinary differential equation in which the component processes are characterized by power-law functions:

$$\frac{dx_i}{dt} = \alpha_i \prod_{j=1}^n x_j^{g_{ij}} - \beta_i \prod_{j=1}^n x_j^{h_{ij}} \quad (i = 1, 2, \dots, n) \quad (1)$$

where n is the total number of state variables or reactants (x_i), i, j ($1 \leq i, j \leq n$) are suffixes of state variables. The terms g_{ij} and h_{ij} are interactive effectivity of x_j to x_i . The first term represents all influences that increase x_i , whereas the second term represents all influences that decrease x_i . In a genetic network context, the non-negative parameters α_i and β_i are called relative inflow and outflow of gene x_i , respectively, and real-valued exponents g_{ij} and h_{ij} are referred to the interrelated coefficients between genes x_j and x_i . Given the experimentally observed time-course data of x_i , the task is to estimate the values of interrelated coefficients g_{ij} , h_{ij} and non-negative constant α_i and β_i , which can be realized experimentally observed time-course of x_i . The S-system formalism has a major disadvantage in that this formalism includes large number of parameters that must be estimated (α_i , β_i , g_{ij} , h_{ij}); the number of estimated parameters in S-system formalism is $2n(n+1)$, where n is the number of state variables (x_i); the number of estimated parameters increases with the second power of the number of system component (n).

2.2. Optimization Procedure

Because the S-system is a formalism of ordinary nonlinear differential equations, the system can easily be solved numerically by using a numerical calculation program to be customized specifically for this structure. However, when an adequate time-course of relevant state variables is given, the set of parameter values α_i , β_i , g_{ij} and h_{ij} , in many cases, will not be uniquely determined, as it is highly possible that other sets of parameter values will also show a similar time-course. Therefore, even if one set of parameter values that explains the observed time-course is obtained, this set is still one of the best candidates to explain the observed time-courses. Our strategy is to explore and exploit these candidates within the immense searching space of parameter values.

In this problem, each set of parameter values to be estimated is evaluated using the following procedure:

Suppose that $x_{d,i,t}^{\text{cal}}$ is the numerically calculated time-course at time t of state variable x_i in the d -th. Data set and that $x_{d,i,t}^{\text{exp}}$ represents the experimentally observed time-course at time t of x_i in the d -th. Data set. The sum of the square values of the relative error between $x_{d,i,t}^{\text{cal}}$

and $x_{d,i,t}^{\text{exp}}$ gives the total relative error E ;

$$E = \sum_{d=1}^D \sum_{i=1}^N \sum_{t=1}^T \left(\frac{x_{d,i,t}^{\text{exp}} - x_{d,i,t}^{\text{cal}}}{x_{d,i,t}^{\text{exp}}} \right)^2 \quad (2)$$

where D is the total number of data sets observed under different experimental conditions, such as gene disruption and over expression, N is the number of experimentally observable state variables, and T is the number of sampling points over time in one experimental condition. The computational task is to determine the set of parameter values that minimizes the objective function E .

2.3. Real-Coded Genetic Algorithms (RCGAs)

The genetic algorithm (GA) is stochastic search and optimization technique based on the mechanism of biological evolution, such as natural selection and natural genetics. The GA can seek out the “best” solution which will be found in regions of the search space containing a relatively high proportion of “good” solutions, that is, the GA can escape from trapping in local minima. The GAs employing real-valued vectors for representation of the chromosomes is called RCGAs [3]. The RCGAs are one of promising evolutionary algorithms and are applied to several real world problems and have shown good results [12,13]. We have developed an efficient computational technique based on the RCGA called UNDX + MGG [14-16] and have applied to the inference of genetic interaction. In this study, we propose a new method using the RCGA, the combination of AREX (adaptive real-coded ensemble crossover) with JGG (just generation gap).

2.3.1. UNDX

The UNDX [4,5], unimodal normal distribution crossover, generates offspring using the normal distribution defined by three parents, as shown in **Figure 1**. Offsprings are generated around the line segment, which is connecting two parents, P_1 and P_2 . The third parent, P_3 ,

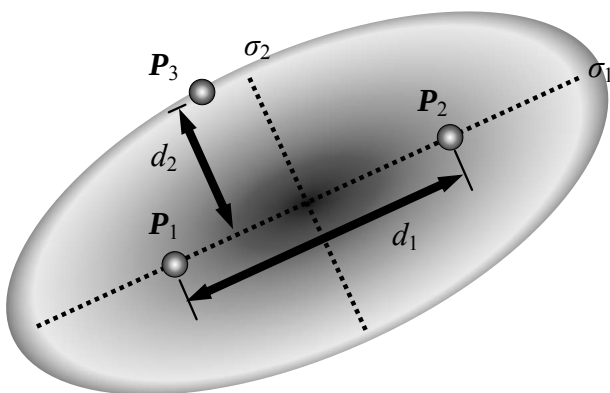


Figure 1. UNDX for parameter search.

is used to determine the standard deviation of the normal distribution. The standard deviation, which corresponds to the coordinate axis along the line segment, is proportional to the distance between P_1 and P_2 . The others are proportional to the distance of P_3 from the line segment and multiplied by $1/\sqrt{l}$, where l is the number of parameters must be estimated; $2n(n+1)$ for S-system. The effect of $1/\sqrt{l}$ is to keep the distribution of offspring around line segment against increasing the number of parameters must be estimated. The mathematical representation of

$$\begin{aligned} c_1 &= m + z_1 e_1 + \sum_{k=2}^l z_k e_k, c_2 = m - z_1 e_1 - \sum_{k=2}^l z_k e_k \\ m &= P_1 + P_2 / 2 \\ z_1 &\sim N(0, \sigma_1^2), z_k \sim N(0, \sigma_2^2) (k=2, 3, \dots, l) \\ \sigma_1 &= \alpha d_1, \sigma_2 = \beta d_2 / \sqrt{l} \\ e_1 &= P_1 + P_2 / |P_1 - P_2| \\ e_i &\perp e_j (i \neq j) (i, j=1, 2, \dots, l) \end{aligned} \quad (3)$$

UNDX is as follows:

where c_1 and c_2 are offspring vector, P_1 and P_2 are parent vectors, m is the middle point of P_1 and P_2 , d_1 is the distance between P_1 and P_2 , d_2 is the distance of the third parent, P_3 , from the line connecting P_1 to P_2 . α and β are constants given by the user, and $\alpha = 0.5$ and $\beta = 0.35$ are recommended. The e_1 is the basis vector in a direction parallel to the coordinate axis along the line segment, and vectors e_i ($i=2, 3, \dots, l$), which are linearly independent basis vectors perpendicular to the vector e_1 , are defined by Gram-Schmidt procedure for finding orthogonal vectors.

2.3.2. MGG

The MGG [6,7] is the generation-alternation model. **Figure 2** shows the conceptual figure of the MGG the procedure of which can be summarized as follows:

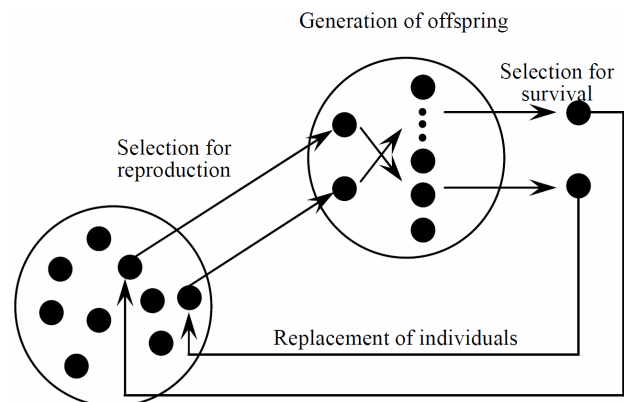


Figure 2. Conceptual figure of minimal generation gap (MGG).

1) Generation of initial population

Make an initial population that is composed of random real number vectors.

2) Selection for reproduction

Select a pair of individuals by random sampling without their placement from the population. The selected pair of individuals becomes the parents of offspring.

3) Generation of offspring

Generate offspring by applying crossover to the parents and evaluate the offspring.

4) Selection for survival

Select two individuals from the family containing the parents and their offspring; one is the best individual and the other is chosen using a roulette-wheel selection. Replace the parents chosen in step 2 in the population with the two selected individuals.

5) Repeat the above procedures from step 2 to step 4 until a certain stop condition is satisfied.

2.3.3. AREX

The AREX [11], adaptive real-coded crossover, is a crossover has been developed to improve the premature convergence in early stage. Akimoto *et al.* has proposed the AREX based on the REX [10,17], real-coded ensemble crossover, which is the generalization of UNDX-m [18]. The AREX generates λ candidate points c_i ($1 \leq i \leq \mu$; μ is the number of parents) around the center $P = (1/\mu) \sum_{j=1}^{\mu} P_j$ of μ parents P_j ($1 \leq i \leq \mu$), as shown in **Figure 3**. And the mathematical representation of the REX is as follows:

$$c_i = P + \alpha \sum_{j=1}^{\mu} \epsilon_{i,j} (P_j - P) \quad (4)$$

where α is the expansion rate, all $\epsilon_{i,j}$ are independently and identically distributed (i.i.d.) random numbers with the average 0 and the variance σ^2 . There is arbitrariness in choice of the probability density function (p.d.f.) of ϵ . For example, if ϵ obeys normal distribution $N(0, \sigma^2)$, the candidate solution generated by Equation (4) obeys

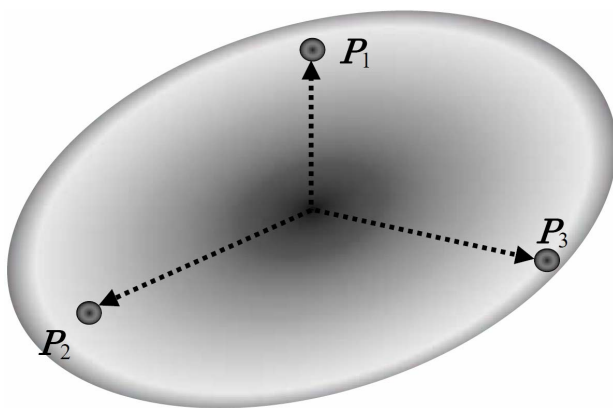


Figure 3. REX using normal distribution.

the normal distribution $N(P, \Sigma)$ with mean vector $\mathbf{0}$ and covariance matrix Equation (5).

$$\Sigma = \alpha^2 \sigma^2 \sum_{j=1}^{\mu} (P_j - P)(P_j - P)^T \quad (5)$$

The REX shows the good performance in ill-conditioned and variable dependent problems because of the invariance against scale transformations and rotation transformations. However, Akimoto *et al.* have reported that there are the situations where the REX is likely to cause early convergence [11]. Furthermore, being combined the adaption of expansion rate (AER) technique and the crossover mean descent (CMD) technique with the REX, they have proposed the AREX to solve the early convergence.

The AER is the expansion rate which is controlled in the middle of the ongoing search process in AREX [11,17]. The CMD is a technique to shift the center of the crossover to promising offspring point. The offspring are generated around the weighted average based on the ranking of evaluation values in parents. The CMD, m , is as follows:

$$m = \sum_{j=1}^{\mu} w_j P_{i,j} \quad (6)$$

where $w_j = 2(\mu + 1 - j) / \mu(\mu + 1)$ is the weight coefficient under the condition of $\sum_{j=1}^{\mu} w_j = 1$.

The AREX is the crossover applying the CMD and the AER to the REX. The mathematical representation of the AREX is as follows:

$$c_i = m + \alpha \sum_{j=1}^{\mu} \epsilon_{i,j} (P_j - P) \quad (7)$$

2.3.4. JGG

The JGG was proposed for multi-parental crossovers [9,10], and it is a modification model of MGG in two major points. First, the JGG does not employ elitist selection in evolution strategies. This modification makes a good effect on multimodal problems. Second, the JGG replaces parents with children completely in every generation. **Figure 4** shows the conceptual figure of the JGG, and the procedure of the JGG can be summarized as follows:

1) Generation of initial population

Make an initial population that is composed of random real number vectors.

2) Selection for reproduction

Select randomly μ parents from the population. The selected μ of individuals becomes the parents of offspring and $\mu = l + 1$ is recommended [9,10], where l is the number of parameters must be estimated.

3) Generation of offspring

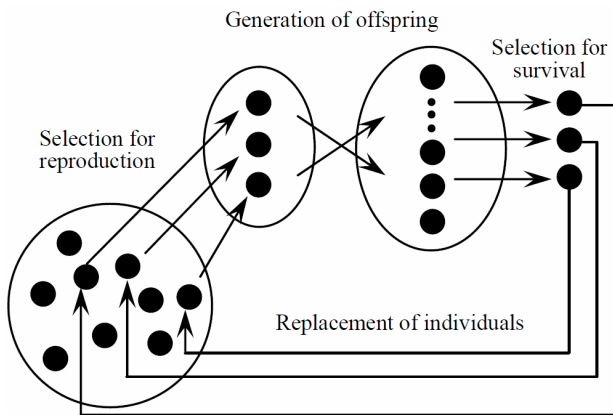


Figure 4. Conceptual figure of just generation gap (JGG).

Generate offspring by applying crossover to the parents and evaluate the offspring.

4) Selection for survival

Select the top ranked μ individuals with respect to the fitness value from the offspring. And replace the parents chosen in step 2 in the population with the μ selected individuals.

5) Repeat the above procedures from step 2 to step 4 until a certain stop condition is satisfied.

3. Results

We prepared an S-system network models composed of six genes, as shown in **Figure 5**, and of seven genes, as shown in **Figure 6**. In these figures, the arrow and T bar represent active and inhibitory interactions, respectively, and the numerals show the value of g_{ij} in S-system formalism. The S-system representation of **Figures 5** and **6** are shown in **Table 1**. We applied the conventional RCGA method (UNDX + MGG) and the proposed method (AREX + JGG) for inferring gene expression network candidates to these artificial gene regulatory network models. We prepared 7 types of time-course data (one a “wild-type” and six “one gene-disrupted strain”) for six genes network model (**Figure 5**) and 8 types of time-course data (one a “wild-type” and seven “one gene-disrupted strain”) for seven genes network model (**Figure 6**), respectively. The number of sampling points in each time-course data set was 70. We calculated time-course data sets for the “one gene-disrupted strain” with the rate constant for the synthetic process of disrupted gene $i(\alpha_i)$ set to 0.

Using these networks, we prepared 4 case studies to compare the performance of UNDX + MGG and AREX + JGG. The case studies 1 to 3 are for the estimation of the parameters of six genes network model, and case study 4 is for the estimation of the parameters of seven genes network model. In each case, the parameters must be estimate dare shown in **Table 1**.

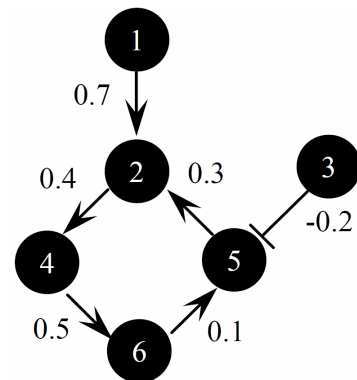


Figure 5. Six-genes network model.

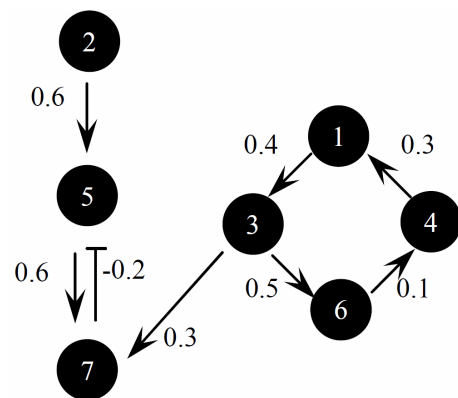


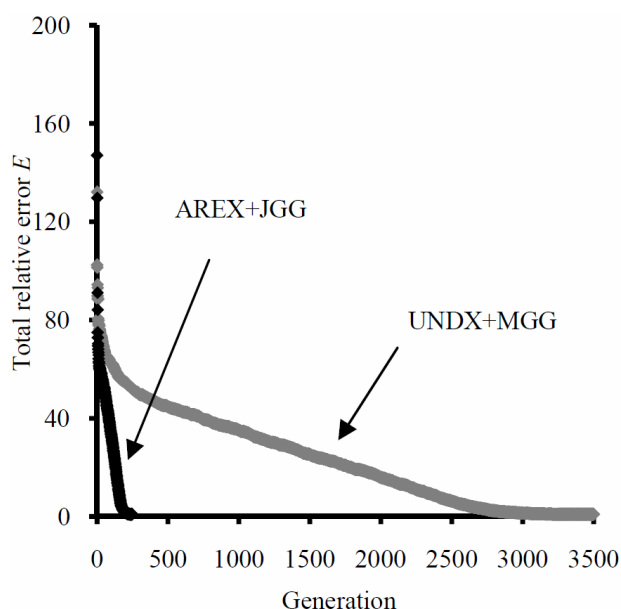
Figure 6. Seven-genes network model.

3.1. Case Study 1

First, in order to compare the convergence speed of two RCGA methods (UNDX + MGG, AREX + JGG), we performed the estimation of parameters in case study 1. In this experiment, we used the recommended default parameter [11,17] of RCGAs as follows: in UNDX + MGG, population size n_{pop} was set to 300, the number of offspring n_c was set to 100, the number of parents n_p was set to 2, and in AREX + JGG, n_{pop} was set to 200, n_c was set to 80, n_p was set to 21. We regarded the optimization terminated successfully when the total relation error E in Equation (2) is less than 0.1. We performed 50 trials, in each RCGA, and evaluated the performances with the average of 50 trials. **Figure 7** shows the average of 50 trials in E of each generation. As shown in **Figure 7**, it is obvious that the convergence speed of AREX + JGG is faster than UNDX + MGG. In all trials, both methods can be estimated the correct values of parameters shown in **Figure 5**. Comparing the CPU time (CPU: XeonE5540 2.53 GHz, Memory: 8.0 GB), AREX + JGG was approximately 20-times faster than UNDX + MGG; the CPU time for AREX + JGG was 38.6 sec and that for UNDX + MGG was 765.0 sec. This experiment shows that AREX + JGG is remarkably better than UNDX + MGG in the convergence speed.

Table 1. Default parameters setting for UNDX + MGG and AREX + JGG.

		Total number of Parameters (l)	S-system parameter must be estimated	Given S-system parameters
Six-gene network model	Case study 1	20	g_{ij} ($i = 2, 4, 5, 6, j = 1, 2, \dots, 6$; when $i \neq j$)	$g_{ij} = 0.0$ (when $i = j$), $h_{ij} = 1.0$ (when $i = j$), $h_{ij} = 0.0$ (when $i \neq j$), $\alpha_i = 1.0$, $\beta_i = 1.0$
	Case study 2	54	g_{ij} ($i = 1, 2, \dots, 6, j = 1, 2, \dots, 6$), h_{ij} ($i = 1, 2, \dots, 6$), α_i ($i = 1, 2, \dots, 6$), β_i ($i = 1, 2, \dots, 6$)	$h_{ij} = 0.0$ (when $i \neq j$)
	Case study 3	84	g_{ij} ($i = 1, 2, \dots, 6, j = 1, 2, \dots, 6$), h_{ij} ($i = 1, 2, \dots, 6, j = 1, 2, \dots, 6$), α_i ($i = 1, 2, \dots, 6$), β_i ($i = 1, 2, \dots, 6$)	None
Seven-gene network model	Case study 4	112	g_{ij} ($i = 1, 2, \dots, 7, j = 1, 2, \dots, 7$), h_{ij} ($i = 1, 2, \dots, 7, j = 1, 2, \dots, 7$), α_i ($i = 1, 2, \dots, 7$), β_i ($i = 1, 2, \dots, 7$)	None

**Figure 7. Convergence profile for AREX + JGG and UNDX + MGG.**

3.2. Case Study 2, 3, 4

Next, in order to evaluate the scalability of optimization, the number of parameters to be estimated increased in case studies 2 to 4. The default parameter of n_c and n_p were set to the recommended values [11] for AREX + JGG. In case studies 2 to 3, in order to find an adequate the default parameter of n_{pop} for AREX + JGG, n_{pop} was scanned between $20l$ and $50l$ (l is the number of estimated parameters). In all runs, the maximum number of generation was set to 100,000. We regarded the optimization terminated successfully when the total relation error E in Equation (2) is less than 0.5. **Table 2** shows the result of convergence speed in UNDX + MGG and AREX + JGG. As shown in **Table 2**, UNDX + MGG could not estimate 54 and 84 parameters within the

maximum limit of generation (100,000). On the other hand, AREX + JGG succeeded the estimation of all parameters except when n_{pop} was set to $20l$. Especially, AREX + JGG showed the best performance when n_{pop} was set to $30l$.

4. Discussion

In this study, we proposed new inferred engine for estimating the gene regulatory network using time-courses of gene expression data using real-coded genetic algorithm (RCGA) with combination of the adaptive real-coded ensemble crossover (AREX) and the just generation gap (JGG). The combination of unimodal normal distribution crossover (UNDX) with the minimal generation gap MGG has disadvantage in convergence and can estimate only a few parameters. The AREX works well in ill-conditioned and variable dependent problems, because the AREX has the invariance against scale transformations and rotation transformations [11]. The JGG is suited for multi-parental crossovers. The JGG is a modification of the MGG in the points that replaces parents with offspring completely every generation and does not employ elitist selection. Akimoto *et al.* reported the AREX + JGG outperforms the existing RCGAs in a number of function evaluations on various functions including functions whose landscape forms ridge structure or multi-peak structure. In this study, AREX + JGG showed remarkably better performance (convergence speed) than UNDX + MGG in all experiments. **Table 2** shows that AREX + JGG could solve the problems in which UNDX + MGG could not do that, and suggests that the adequate default parameter of the population size for AREX + JGG is $30l$, where l is the total number of parameters. By employing AREX + JGG, we succeeded in improving the performances in the point of the convergence speed and the scalability in the number of estimated parameters; compared with UNDX + MGG, the

Table 2. Results of convergence speed in UNDX + MGG and AREX + JGG.

	Total number of Parameters (l)	Given parameters			Total relative error (E)	Last generation	CPU time (sec)/run	
		n_{pop}	n_c	n_p				
UNDX + MGG	54 (Case study 2)	300		100	2	0.83	>100,000	27885.2
		1080	20/	216	2	0.89	>100,000	57680.5
		1620	30/	216	2	1.34	>100,000	74510.4
		2160	40/	216	2	4.00	>100,000	53628.4
	84 (Case study 3)	300		100	2	2.46	>100,000	27033.0
		1380	20/	336	2	3.37	>100,000	113824.0
		2520	30/	336	2	4.58	>100,000	105570.0
		3360	40/	336	2	5.73	>100,000	114232.2
	54 (Case study 2)	1080	20/	216	55	0.67	>100,000	59190.8
		1620	30/	216	55	0.50	2374	1367.2
2160		40/	216	55	0.50	3759	2177.3	
2700		50/	216	55	0.50	3871	2228.2	
AREX + JGG	84 (Case study 3)	1380	20/	336	85	0.70	>100,000	139688.0
		2520	30/	336	85	0.50	3335	2801.6
	3360	40/	336	85	0.50	3931	3511.8	
	4200	50/	336	85	0.50	4682	4214.7	
	112 (Case study 4)	3360	30/	448	113	0.50	4388	7449.0

convergence speed and the scalability made rapid progress to approximately 5-fold and 20-fold, respectively.

As described already, in S-system formalism, the number of parameters to be estimated is proportion to the second power of the number of system components (n). In this study, AREX + JGG could estimate maximum 112 parameters, which corresponds to $n = 7$. Suppose we apply this method to the inference of regulatory network of all amino acids, we have to estimate 840 parameters because the total number of amino acids is 20 ($n = 20$). For the larger network estimation, we should further improve the proposed method. Substantially the GA can find the solution within a comparatively large searching range, but it is stepwise convergence, because the GA does not have an efficient local searching function. One of the methods having a local searching function is the modified Powell method. The modified Powell method is well known to have an ultimate fast convergence among the various direct-search methods without calculating the derivative of the objective function, especially where the objective function is well approximated by a quadratic form of parameters to be estimated. The modified Powell method can find the solution very quickly only when the initial guess is very near the solution; this method has no way to escape from a local minimum. The hybrid method, the RCGA and the modified Powell method, is expected to offer all advantages of both optimization techniques. We are now on going the development of the hybrid

method by incorporation the modified Powell method into AREX + JGG, which is expected to become a more powerful optimization method for the larger network estimation.

5. Acknowledgements

This study was supposed by Grant-in-Aid for Scientific Research on Innovative Areas (Research in a proposed area), "Synthetic Biology for the Comprehension of Biomolecular Networks" from the Ministry of Education, Culture, Sports, Science and Technology, Japan (No. 23119001 (M. Okamoto)).

REFERENCES

- [1] M. A. Savageau, "Biochemical Systems Analysis: A Study of Function and Design in Molecular Biology," Addison-Wesley, Reading, Boston, 1976.
- [2] D. Tominaga, N. Koga and M. Okamoto, "Efficient Numerical Optimization Algorithm Based on Genetic Algorithm for Inverse Problem," *Proceedings of the Genetic and Evolutionary Computation Conference*, Las Vegas, 8-12 July 2000, p. 251.
- [3] L. J. Eshleman and J. D. Schaffer, "Real-Coded Genetic Algorithms and Interval-Schemata," *Foundations of Genetic Algorithms*, Vol. 2, 1993, pp. 187-202.
- [4] I. Ono and S. Kobayashi, "A Real-Coded Genetic Algorithm for Function Optimization Using Unimodal Normal

- Distribution Crossover,” *Journal of Japanese Society for Artificial Intelligence*, Vol. 14, No. 6, 1999, pp. 1146-1155.
- [5] I. Ono, S. Kobayashi and K. Yoshida, “Global and Multi-Objective Optimization for Lens Design by Real-Coded Genetic Algorithms,” *International Optical Design Conference Proceedings of SPIE*, Vol. 3482, 1998, pp. 110-121.
- [6] H. Sato, I. Ono and S. Kobayashi, “A New Generation Alternation Model of Genetic Algorithms and Its Assessment,” *Journal of Japanese Society for Artificial Intelligence*, Vol. 12, No. 5, 1997, pp. 734-744.
- [7] H. Satoh, M. Yamamura and S. Kobayashi, “Minimal Generation Gap Model for GAs Considering Both Exploration and Exploitation,” *Proceedings of 4th International Conference on Soft Computing*, Iizuka, 30 September-5 October 1996, pp. 494-497.
- [8] N. Shikata, Y. Maki, M. Nakatsui, M. Mori, Y. Noguchi, S. Yoshida, M. Takahashi, N. Kondo and M. Okamoto, “Determining Important Regulatory Relations of Amino Acids from Dynamic Network Analysis of Plasma Amino Acids,” *Amino Acids*, Vol. 38, No. 1, 2009, pp. 179-187. [doi:10.1007/s00726-008-0226-3](https://doi.org/10.1007/s00726-008-0226-3)
- [9] Y. Akimoto, R. Hasada, J. Sakuma, I. Ono and S. Kobayashi, “Generation Alternation Model for Real-Coded GA Using Multi-parent: Proposal and Evaluation of Just Generation Gap (JGG),” *Proceedings of the 19th SICE Symposium on Decentralized Autonomous Systems*, Tokyo, 29-30 January 2007, pp. 341-346.
- [10] S. Kobayashi, “The Frontiers of Real-Coded Genetic Algorithms,” *Journal of Japanese Society for Artificial Intelligence*, Vol. 24, No. 1, 2009, pp. 147-162.
- [11] Y. Akimoto, Y. Nagata, J. Sakuma, I. Ono and S. Kobayashi, “Proposal and Evaluation of Adaptive Real-Coded Crossover AREX,” *Journal of Japanese Society for Artificial Intelligence*, Vol. 24, No. 6, 2009, pp. 446-458.
- [12] J. Sakuma and S. Kobayashi, “Latent Variable Crossover Fork-Tablet Structures and Its Application to Lens Design Problems,” *Proceedings of the Genetic and Evolutionary Computation Conference (GECCO-2005)*, Washington DC, 2005, pp. 1347-1353.
- [13] I. Ono, H. Kita and S. Kobayashi, “A Robust Real-Coded Genetic Algorithm Using Unimodal Normal Distribution Crossover Augmented by Uniform Crossover: Effects of Self-Adaptation of Crossover Probabilities,” *Proceedings of the Genetic and Evolutionary Computation Conference (GECCO-1999)*, Orlando, 1999, pp. 496-503.
- [14] T. Ueda, N. Koga, I. Ono and M. Okamoto, “Efficient-Numerical Optimization Technique Based on Real-Coded Genetic Algorithm for Inverse Problem,” *Proceedings of 7th International Symposium on Artificial Life and Robotics*, Beppu, Oita, 16-18 January 2002, pp. 290-293.
- [15] T. Ueda, N. Koga, I. Ono and M. Okamoto, “Development of Efficient Numerical Optimization Method Based on Real-Coded Genetic Algorithm: Application to the Estimation of Large Number of Real-Valued Parameters,” *Proceedings of the 7th International Symposium for Biochemical Systems Theory: From Phenotype to Genotype and Back*, Averoy, More og Romsdal, 17-20 June 2002, pp. 43-44.
- [16] M. Nakatsui, T. Ueda, Y. Maki, I. Ono and M. Okamoto, “Method for Inferring and Extracting Reliable Genetic Interactions from Time-Series Profile of Gene Expression,” *Mathematical Biosciences*, Vol. 215, No. 1, 2008, pp. 105-114. [doi:10.1016/j.mbs.2008.06.007](https://doi.org/10.1016/j.mbs.2008.06.007)
- [17] Y. Akimoto, J. Sakuma, I. Ono and S. Kobayashi, “Adaptation of Expansion Rate for Real-Coded Crossovers,” *Proceedings of the Genetic and Evolutionary Computation Conference (GECCO-2009)*, Montreal, 8-12 July 2009, pp. 739-746.
- [18] H. Kita, I. Ono and S. Kobayashi, “Multi-Parental Extension of the Unimodal Normal Distribution Crossover for Real-Coded Genetic Algorithms,” *Proceedings of the IEEE Congress on Evolutionary Computation*, Washington DC, 6-9 July 1999, pp. 1581-1587.