A Constructive Genetic Algorithm For The Generalized Assignment Problem

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Abstract

We present in this paper an application of the *Constructive Genetic Algorithm (CGA)* to the *Generalized Assignment Problem (GAP)*. The *CGA* presents some new features compared to a traditional *genetic algorithm (GA)*, such as a population formed only by schemata, recombination among schemata, dynamic population, mutation in structures, and the possibility of using heuristics in schemata and/or structure representation. The *GAP* can be described as a problem of assigning *n* items to *m* knapsacks, n > m, such that each item is assigned to exactly one knapsack, subject to capacity constraints on the knapsacks. In our application of *CGA to GAP*, we regard the *GAP* as a clustering problem. A binary representation is used for schemata and structures, and an assignment heuristic allocates items to knapsacks. Schemata do not consider all the problem data. The schemata are recombined, and they can produce new schemata or structures. New schemata are evaluated and can be added to the population if they pass an evolution test. Structures can result from recombination of schemata or complementing of good schemata. They suffer mutation and the best structure generated is kept in the process. Computational tests have been performed using instances of large scale available in the literature.

<u>1.</u> Introduction

Genetic Algorithms (GA) have become popular in recent years as efficient heuristics for difficult combinatorial optimisation problems. The underlying foundation for such algorithms are the controlled evolution of a structured population. Today there are many variations on the general GA theme and all such variations can be classified generically as population heuristics [3], that is as heuristics that operate with a population of solutions. Such heuristics are in marked contrast to other approaches, such as tabu search and simulated annealing, that operate on just a single solution.

The *GA* works on a set of variables called *structures*. For application to optimization problems [4,14], the first step is the definition of a codification plan that allows a one to one mapping between solutions and structures. The following string can represent a structure $S_k = (s_{k1}, s_{k2}, ..., s_{kn})$, where *n* is the number of variables in the problem. A *fitness function* assigns a numeric value to each member of the current *population* (a collection of structures). The *genetic operators* used are *selection*, like tournament or biased roulette wheel, working together with a number of *crossover* and *mutation* operators. The best structure is kept after a predefined number of generations [10,17,18].

Holland [10] put forward the *building block* hypothesis (*schema* formation and conservation) as a theoretical basis for the *GA* mechanism. In his view avoiding disruption of good schema is the basis for the good behaviour of a *GA*. One major problem with building blocks however is that schemata are evaluated indirectly, via evaluation of their instances (structures), rather than directly, as an instance may typically represent several schemata. The *Constructive Genetic Algorithm* (*CGA*) [13] was

proposed recently as an alternative to the traditional *GA* approach [9,20]. One of the objectives of a *CGA* is the direct evaluation of schemata.

Some steps in *CGA* are notably different from a classical *GA*. Problems are modeled as *bi-objective* optimization problems, which consider the evaluation of two fitness functions. The evolution process is conducted to attain the two objectives conserving schemata that survive to an adaptive threshold test. Consequently the *CGA* works with a *dynamic population*, composed exclusively of schemata, which increases after the use of *recombination operators* and can decrease as generations pass, guided by an *evolution parameter*. Schemata recombination diversifies the population thereby generating new schemata or structures. A structure can be obtained after schemata recombination, or by complementing a good schema. They suffer mutation and are compared to the best one found so far, which is always retained. Another main difference between a classical *GA* and a *CGA* is the new *fg-fitness* process (see [13] for further details).

We examine in this paper a CGA application to the problem of the minimum cost assignment of *n* tasks to *m* agents (n > m), such that each task is assigned to only one agent subject to capacity constraints on the agents. This problem is an important combinatorial optimization problem, the *Generalized Assignment Problem (GAP)*.

Many real life applications can be modeled as a *GAP*, e.g. resource scheduling, the allocation of memory space in a computer, the design of a communication network with capacity constraints for each network node, assigning software development tasks to programmers, assigning jobs to computers in a network, vehicle routing problems, and

GAP is NP-hard [19]. A number of algorithms in the literature are exact tree search methods [16,21], and there are also a number of heuristics for the problem [4,8,11,12,15,16,19].

This paper is organized as follow. The *CGA* application to the *GAP* is presented in Section 2. The *GAP* is modeled as a bi-objective optimization problem that drives the evolutionary search for well adapted structures (solutions) and good schemata. The relevant aspects of the *CGA* are explained, the schemata and structure representation, the evolution process, selection, recombination and mutation, and a *CGA* pseudo-code. Section 3 presents computational tests considering large scale instances from the literature, providing insights into *CGA* performance.

2. CGA modeling

The *CGA* must be tailored to be applied to the *GAP*. A typical mathematical formulation to the *GAP* is presented in the following.

The *GAP* is best described using knapsack problems [16]. Given *n* items and *m* knapsacks, with p_{ij} as the cost associated with assigning item *j* to knapsack *i*, w_{ij} as the weight of assigning item *j* to knapsack *i*, and c_i the capacity of knapsack *i*, assign each item *j* to exactly one knapsack *i*, not exceeding knapsack capacities. Then the *GAP* can be formulated as

$$v(GAP) = Min \sum_{i=1}^{m} \sum_{j=1}^{n} p_{ij} x_{ij}$$
(GAP) subject to $\sum_{j=1}^{n} w_{ij} x_{ij} \le c_i, i \in M = \{1, ..., m\},$

$$\sum_{i=1}^{m} x_{ij} = 1, j \in N = \{1, ..., n\},$$

$$x_{ij} \in \{0, 1\}, i \in M, j \in N.$$

For the *CGA* application, the *GAP* will be formulated as a bi-objective optimization problem. While in the other evolutionary algorithms, the evaluation of individuals is based on a single function (the fitness function), in *CGA* this process relies on two functions, mapping the space of schemata onto \Re_+ . We first describe the structure and schema representation.

2.1. Schema and structure representation

For schema and structure representation, we used a sequence of n symbols, where n is the

number of items. Seed items are initially assigned to the *m* knapsacks, exactly one per knapsack. If the structure is a schema, some items are not considered, i.e. they are considered temporarily out of the problem. The structures have in each position one of the following three possible symbols:

- 1 to indicate the seed item is assigned to a knapsack,
- 0 to indicate a non-seed item assigned to a knapsack, and
- # to indicate items temporarily out of the problem.

A structure with #'s represents a schema. For example considering a problem with 10 items and 3 knapsacks, a structure could be represented by $S_k = (\#,1,\#,0,1,0,0,\#,1,0)$, where item number 2 was assigned to knapsack 1, item number 5 was assigned to knapsack 2, and item number 9 is assigned to knapsack 3. The items receiving labels 0 will be assigned to one of the knapsacks according to an assignment heuristic, and the items with labels # are out of the problem.

Suppose we have a given schema or structure S_k . The following assignment heuristic is used to complement the S_k representation:

Assignment Heuristic - AH

- 3 Assigning the other items to the knapsacks (labels 0 and #)
 - 3.1 solve the m knapsack problems separately exactly
 - 3.2 update the knapsack capacities for the items assigned to exactly one knapsack,
 - 3.3 resolve the m knapsack problems separately exactly for the remaining items,
 - 3.4-update the knapsack capacities for the items assigned to exactly one knapsack,
 - 3.5 for each item j remaining, assign it to knapsack_{i*} corresponding to the smallest w_{i*j} .
 - 3.5 If the obtained solution is not feasible to GAP, restart the assignments of the n-m items (the

^{1 -} Assign the m items with label 1 to the m knapsacks,

^{2 –} Update the knapsack capacities,

m seed items were already assigned in step 1), assigning item *j* to knapsack_{*i**} corresponding to the smallest w_{i*j} . If capacities are violated, assign if possible, item *j* to the knapsack corresponding to the next smallest w_{ij} for which capacities are not violated.

4 - If the solution is feasible to GAP improve the solution with the second part of MMTH (see [15]), else discard the schema and select a new one.

5 – Discard from knapsacks the items with labels # in S_k .

Knapsack problems are solved exactly using the algorithm of Horowitz and Sahni [16].

2.2. The bi-objective problem

Let C be the set of all structures and schemata that can be generated by the 0-1-# string representation of section 2.1., and consider two functions f and g, defined as $f : C \circledast$ \Re_+ and $g : C \circledast \Re_+$ such that $f(S_k) \pounds g(S_k)$, for all $S_k \in C$. We define the double fitness evaluation of a structure or schema s_i , due to functions f and g, as fg-fitness.

The CGA optimization problem implements the *fg-fitness* considering two objectives:

- (*interval minimization*) Search for $S_k \in C$ of minimal $\{g(S_k) f(S_k)\}$, and
- (*g* maximization) Search for $S_k \in C$ of maximal $g(S_k)$.

Considering the schema representation, the fg-fitness evaluation increases as the number of labels # decreases and therefore structures or schemata with few labels # have higher fg-fitness.

To attain these purposes, a problem to be solved using *CGA* is modeled as the following *Bi-objective Optimization Problems (BOP)*:

$$Min \quad \{g(S_k) - f(S_k)\}$$
$$Max \quad g(S_k)$$
$$subj. \text{ to } \quad g(S_k) \ge f(S_k)$$
$$S_k \in \mathbf{C}$$

Functions f and g must be properly identified to represent optimization objective of the *GAP*. The application of the *CGA* to the *GAP* is made through an analogy with clustering problems. Each knapsack is a capacitated cluster to which items must be allocated

Consider a structure or schema $S_k \in X$. For the *GAP*, after the application of the assignment heuristic *AH*, the clusters $C_i(S_k)_{AH}$ are identified, corresponding to index of

items on knapsack_i, i=1,...,m. The function g is then defined by $g(S_k) = \sum_{i=1}^m \sum_{j \in C_i(S_k)_{AH}} p_{ij}$.

To define the function f the following *MAH* heuristic is applied to S_k , producing an additional move of one item between two knapsacks:

Modified Assignment Heuristic – MAH

After the MAH application, if S_k is an structure, the corresponding GAP solution may be

^{1.} Apply AH to S_k .

^{2.} Sort in non-increasing order the costs p_{ij} corresponding to the items in knapsacks presenting label 0 in S_k .

^{3.} Let p_{i*j*} be the cost of the item at the first order position (item j^* was assigned to knapsack i^*).

^{4.} Sort in non-decreasing order the costs p_{ij*} , i=1,...,m. Let $p_{i'j*}$ be the cost of the item at the first order position.

^{5.} Move item j^* to knapsack i'.

infeasible. The new clusters $C_i(S_k)_{MAH}$ are used in the definition of function f as

$$f(S_k) = \sum_{i=1}^{m} \sum_{j \in C_i(S_k)_{MAH}} p_{ij}$$
. Clearly we have that $f(S_k) \pounds g(S_k)$

The interval $g(S_k) - f(S_k)$ can be interpreted as the cost of a wrong assignment if the resulting *GAP* solution is still feasible, but in general it tries to reduce de overall assigned costs p_{ij} .

2.3. The evolution process

The *BOP* defined in *section 2.2* is not directly considered as the set X is not completely known. Alternatively we consider an evolution process to attain the objectives (*interval minimization and g maximization*) of the *BOP*.

At the beginning of the process, two *expected values* are given to these objectives, a nonnegative real number $g_{\max} > Max_{S_k \in X} g(S_k)$, that is an upper bound to $g(S_k)$, for each S_k $\hat{I} C$, and the interval length dg_{\max} , obtained from g_{\max} using a real number $0 < d \le 1$.

The evolution process is then conducted considering an adaptive rejection threshold, which contemplates both objectives in *BOP*. Given a parameter a^{30} , the expression

$$g(S_k) - f(S_k)^3 dg_{\max} - \mathbf{a} \cdot d[g_{\max} - g(S_k)]$$
 (2.3.1)

presents a condition for rejection from the current population of a schema S_k .

The right hand side of (2.3.1) is the threshold, composed of the expected value to the interval minimization dg_{max} , and the measure $[g_{max} - g(S_k)]$, that shows the difference of $g(S_k)$ and g_{max} evaluations. For $\mathbf{a} = 0$, (2.3.1) is equivalent to comparing the interval length obtained by S_k and the expected length dg_{max} . Schemata are discarded if expression (2.3.1) is satisfied. When $\mathbf{a} > 0$, schemata with most labels # in representation have higher possibility of being first discarded, as they present, in general, smaller differences $[g_{max} - g(S_k)]$.

Parameter a is related to time in the evolution process. Considering that the good schemata need to be preserved for recombination, the *evolution parameter* a starts from 0, and then increases slowly, in small time intervals, from generation to generation. The population at the evolution time a, denoted by P_a , is dynamic in size according to the value of the adaptive parameter a, and can be emptied during the process.

The parameter \mathbf{a} is now isolated in expression (2.3.1), thus yielding the following expression and corresponding rank to S_k :

$$\boldsymbol{a} \geq \frac{dg_{\max} - [g(S_k) - f(S_k)]}{d[g_{\max} - g(S_k)]} = \boldsymbol{d}(S_k).$$

At the time they are created, schemata receive their corresponding rank value $G(S_k)$. The *rank* of each schema or structure is compared with the current evolution parameter **a**. At the moment a schema is created, it is then possible to have some figure of its survivability. The higher the value of $G(S_k)$, and better is the schema to the *BOP*, and they also have more surviving and recombination time.

2.4. Selection and recombination

Selection of individuals can be made in several ways. *CGA* has been tested with a number of optimization problems and in all cases an appropriate approach is that the population is kept ordered using a key value that considers the *fg-fitness* and its proximity to a feasible solution representation (structure). Then, several times in a generation, two schemata are randomly selected, one from among the best part of the population and the other from the whole population, and these are recombined to form (one or more) new schemata or structures (see Lorena and Furtado [13]).

The recombination is made depending on the problem and the way the structure represents a solution. The main goal of recombination is population diversification. Structures representing feasible solutions can be generated not only by recombination, but also by complementation of a selected schema. The best results found with the *CGA* uses mutation over structures that represent feasible solutions for the problem (see Lorena and Furtado [13]).

The population is then kept non-decreasing ordered according the following key $\Delta(S_k) = \frac{1+d_k}{n-n_{\#}}, \text{ where } d_k = \frac{g(S_k) - f(S_k)}{g(S_k)} \text{ and } n_{\#} \text{ is the number of } \# \text{ labels in } S_k.$

Schemata with small $n_{\#}$ and/or presenting small d_k are better and appear in first order

positions.

The method used for selection takes one schema from the *n* first positions in the population (*base*) and the second schema from the whole population (*guide*). Before recombination, the first schema is complemented to generate a structure representing a feasible solution, i.e. all #'s are replaced by 0's. This complete structure suffers mutation and is compared to the best solution found so far (which is kept throughout the process).

The recombination merges information from both selected schemata, but preserves the number of labels 1 (number of knapsacks) in the new generated schema.

Recombination

$$\begin{split} & if \ S_{base}(j) = S_{guide}(j) \ then \ S_{new}(j) \neg \ S_{base}(j) \\ & if \ S_{guide}(j) = \# \ then \ S_{new}(j) \neg \ S_{base}(j) \\ & if \ S_{base}(j) = \# \ or \ 0 \ and \ S_{guide}(j) = 1 \ then \\ & S_{new}(j) \neg \ 1 \ and \ S_{new}(i) \neg \ 0 \ for \ some \ S_{new}(i) = 1 \\ & if \ S_{base}(j) = 1 \ and \ S_{guide}(j) = 0 \ then \\ & S_{new}(j) \neg \ 0 \ and \ S_{new}(i) \neg \ 1 \ for \ some \ S_{new}(i) = 0 \end{split}$$

At each generation, exactly n new schemata are created by recombination. If a new schema does not represent a feasible solution, then it is inserted into the population; otherwise it suffers mutation and is compared to the best solution found so far. The following pseudo-code describes the mutation process:

The mutation process was limited to considering just ten new structures to avoid excessive computation time.

At each generation, after new schemata insertion, the population is scanned to remove all structures satisfying the condition $\mathbf{a} \ge \mathbf{c}(S_k)$. As described earlier in this paper, the evolution parameter \mathbf{a} is initially set to zero and slowly increased at each generation.

2.5. The CGA pseudo-code

The basic form of a CGA [13] is:

Constructive Genetic Algorithm - CGA

```
\alpha := 0;
\varepsilon := 0.01;
                                                                                       { time interval }
Initialize P_{\alpha};
                                                                                       { initial population }
Evaluate P_{\alpha};
                                                                                       { proportional fitness }
For all S_k \hat{\mathbf{I}} P_{\mathbf{a}} compute \mathbf{G}(S_k)
                                                                                       { rank computation }
end_for
While (not stop condition) do
            For all S_k \mathbf{\hat{I}} P_a satisfying \alpha < \mathbf{\alpha}(S_k) do
                                                                                      { evolution test }
                         \alpha := \alpha + \varepsilon;
                         Select P_{\alpha} from P_{\alpha-\epsilon};
                                                                                       { reproduction operator }
                         Recombine P_{\alpha};
                                                                                       { recombination operators }
                         Evaluate P_{\alpha};
                                                                                       { proportional fitness }
            end_for
            For all new S_k \hat{\mathbf{I}} P_{\mathbf{a}} compute \mathbf{C}(S_k)
                                                                                      { rank computation }
            end_for
end while
```

As the evolution parameter α increases, the population size initially increases and then start to decrease until eventually the population becomes empty. Two stopping conditions are considered: the process stops when the population is empty, or when a pre-defined generation limit is reached.

To compute the upper bound g_{max} , at the very beginning of the process, a structure S representing a feasible solution (no #'s) is randomly generated and g(S) is taken as the g_{max} value.

For all the computational results presented in this paper an initial population was randomly created with 20% of positions in each structure with label 0, exactly m (number of knapsacks) with label 1, and the remaining positions having the label #.

3. Results

In this section we outline the CGA performance on the GAP. The CGA was coded in C and run on a SUN ULTRA SERVER 2, 200 MHz machine.

A set of large-scale instances were solved (of dimensions, m x n, (5 x 100), (5 x 200), (10 x 100), (10 x 200), (20 x 100) and (20 x 200), from OR-Library [2]). These comprise 24 instances of different sizes and types. Referring to *Table 1* the problems in classes *A*, *B* and *C* present increasingly constrained knapsacks. Class *D* comprises more difficult correlated problems.

Table 1 presents the best *CGA* results (best $g(S_k)$) for ten replications compared with the best known solutions reported in [5]. The *CGA* parameters are set to:

d = 0.15, **a** starts at 0, **e** = 0.1 for 0 **£a £1**, **e** = 0.01 for **a** > 1. The stopping conditions: maximum number of generations = 150, or the population is empty (**a** is big enough).

For problems in class A the best known solutions are optimal so the algorithm was terminated when those solutions were found.

The *CGA* solutions reported in *Table 1* are very close to the best known solutions, obtained in the *GA* implementation of Chu and Beasley [5] who ran their *GA* until 500000 distinct feasible solutions were found. It can be conjectured that the computational efforts of *CGA* are very small compared to their *GA*. The computer times are not directly comparable, as the *GA* was run on a different machine.

Problem	Best known	CGA solution	Number of	CGA times
	solution		generations	(seconds)
A 5x100	1698	1698	51	253
A 5x200	3235	3235	1	502
A 10x100	1360	1360	87	308
A 10x200	2623	2623	72	930
A 20x100	1158	1158	1	350
A 20x200	2339	2339	19	860
B 5x100	1843	1843	150	302
B 5x200	3553	3601	150	432
B 10x100	1407	1410	150	165
B 10x200	2831	2831	150	949
B 20x100	1166	1166	150	474
B 20x200	2340	2347	150	683
C 5x100	1931	1941	150	195
C 5x200	3458	3460	150	405
C 10x100	1403	1423	150	203
C 10x200	2814	2815	150	498
C 20x100	1244	1244	150	479
C 20x200	2397	2397	150	1059
D 5x100	6373	6479	150	259
D 5x200	12796	12823	150	1253
D 10x100	6379	6390	150	497
D 10x200	12601	12634	150	1321
D 20x100	6269	6280	150	974
D 20x200	12452	12471	150	2158

Table 1: Computational results

4. Conclusions

In this paper we have presented an application of the constructive genetic algorithm to the generalized assignment problem. Computational results were promising as compared to a previous genetic algorithm approach presented in the literature.

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