Investigation of simulated annealing components to solve the university course timetabling problem

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Abstract. Simulated Annealing (SA) is a common meta-heuristic algorithm that has been widely used to solve complex optimization problems. This is due to its ease of implementation and capability to escape from local optimum. This research conducts an investigation on three of SA components: the initial temperature, cooling schedule and neighborhood structure. We observed that the high initial temperature leads SA to accept any solution (wasting more computational time), whilst the lower value leads SA to quickly trap in local optimum. Based on research findings from this phase, for each component we suggested a technique to overcome the limitations. The limitations are: (i) a dynamic initial temperature mechanism that dynamically chose the suitable initial temperature for each instance problem; (ii) adaptive cooling schedule that will adjust the temperature value during the search; and (iii) a new neighborhood structure that will improve the search ability by minimizing the random selection. In the second phase. The experimental results show that the proposed techniques and approaches in all phases have outperformed the SA and comparable to other approaches in the literature (tested on university course timetabling benchmark dataset ITC2007-Track3).

Keywords: simulated annealing, meta-heuristic, university course timetabling problem, optimization, ITC2007-Track3.

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1. Introduction

Timetabling is considered as a unique case of scheduling problems [23]. Wren [23] defined the timetabling as follows:

“The allocation, subject to constraints, of given resources to objects being placed in spacetime, in such a way as to satisfy as nearly as possible a set of desirable objectives”.

Schaerf [18] categorized the educational field into three common timetabling problems: university course timetabling problem, examination timetabling problem and school timetabling problem. All of them may different but they still have the same fundamental characteristics. For example, the sizes of class for all school lectures are normally similar and have the same group of students in common that are associated with the same courses, meanwhile the university courses have different number of students’ enrolments. Table (1) summarizes some of the common meta-heuristic algorithms related to different categories which are presented by Birattari et al. [4], where √ means that the algorithm is totally categorized under this category and x means it does not, whilst ¬ means it is a partly related to this category.

<table>
<thead>
<tr>
<th>Category</th>
<th>Trajectory</th>
<th>Population</th>
<th>Memory</th>
<th>Various neighborhood</th>
<th>Dynamic</th>
<th>Nature inspired</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SA</td>
<td>TS</td>
<td>GA</td>
<td>ACO</td>
<td>ILS</td>
<td>grasp</td>
</tr>
<tr>
<td>Trajectory</td>
<td>√</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>√</td>
</tr>
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<td>x</td>
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<td>x</td>
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<tr>
<td>Dynamic</td>
<td>x</td>
<td>¬ x</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Nature inspired</td>
<td>√</td>
<td>x</td>
<td></td>
<td>√</td>
<td>x</td>
<td>x</td>
</tr>
</tbody>
</table>

Note: ACO: ant colony optimization; TS: Tabu search; ILS: iterated local search; GRASP: greedy randomized adaptive search procedure; SA: simulated annealing; GLS: guided local search, GA: genetic algorithm.

SA is a trajectory-based algorithm or a single point algorithm [4]. The single point algorithms consider a single element of the solution space at each iteration by jumping from position to another in the solution space.

SA uses a special strategy to escape from local optimum. At first, SA will explore the search space (diversification) by easily accepting worse solution based on probability acceptance criterion. Later on, the probability is decreased during the search space process that leads to focus more on promising regions (intensification) [1]. In fact, the acceptance probability depends on the temperature which is reduced during the search process to give balance between the diversification and intensification. However, SA could be trapped into local optimum and may consume a longer time to find good solutions [24]. Hence, many researchers attempted to improve the SA performance by using different approaches, such as adaptive SA (ASA) suggested by Ingber [10], or by
hybridize SA with other meta-heuristics approaches, such as genetic algorithm [21]. In fact, hybrid techniques can achieve different advantages; such as the best performance; by combining the advantages of each individual technique into one hybrid algorithm. Therefore, this research is motivated to investigate the process of SA performance improvement by escaping from local optimum and reducing the computational time.

This research focuses on improving a SA performance by investigating the effect of SA components: temperature, cooling schedule and neighborhood structure on the SA performance. The good initial temperature will balance the diversification and intensification properly by avoiding the waste of computational time when the initial temperature is high or when trap in local optimum which is a result of a very low temperature [22].

Elmohamed et al. [8] claimed that adjusting the decrement amount of the given temperature during the search process is an important issue, in order to avoid the wasting of computational time. Furthermore, The high temperature with slow cooling schedule leads SA to accept many bad solutions (a zigzag walk), while the low temperature with fast cooling schedule leads SA to trapped quickly in the local optimum by rejecting most of the unimproved solutions and accepting the improved solutions only (descent method). Meanwhile, an effective neighborhood structures may easily improve the solution quality [14]. Moreover, choosing the most suitable neighborhood is an important issue to avoid the discounted neighborhood and save the computational time [15].

AlHadid et al. [2] proposed a hybrid SA with EMC technique to divert the search effectively to another promising region by escaping the search space from local optimum to another promising region space. AlHadid et al. [2] stated that the proposed technique results has outperformed the standard SA and gave comparable results to other approaches when tested on ITC2007-Track3 university course timetabling datasets.

According to the previous dissection, we have identified two main research questions (in regard to SA algorithm) to be answered:

a) How can we enhance the SA performance by escaping from local optimum?

b) How can we reduce the SA computational time?

In order to answer the main research questions, we have identified four research questions which need to be answered, the questions are:

Q1) How to initialize the initial temperature dynamically based on the problem instance individually?

Q2) What is the effective cooling schedule that can adaptively adjust the amounts of temperature decrement during the SA search process?

Q3) How to design a good neighborhood structure that can help SA to effectively search for good quality solution and minimize the random swap and move?

Q4) How to choose a good neighborhood structure among different neighborhood structures during the search process that is able to avoid the disconnected neighborhood?
2. Related work

2.1 Dynamic initial temperature

This section presents Dynamic Initial Temperature (D-IT) which proposed by Tarawneh et al. [20] to improve performance of SA by initializing the initial temperature dynamically. Normally, when the temperatures are very high the search space of SA could be very wide by accepting many worse solutions. Meanwhile, the range of accepting worse solution(s) becomes very small when the initial temperature is low.

The proposed D-IT mechanism by Tarawneh et al. [20] is used to estimate the suitable range of accepting worse solution at early stage, by performing a preliminary experiment to clarify the acceptance ratio related to different worse solutions for each temperature and deviation average $\gamma$, where $\gamma$ is calculated using the following equation:

$$
\gamma = \frac{\sum_{i=1}^{n} |\Delta f|}{n}
$$

Where $\Delta f = f(s^*) - f(s)$ and $n$ is the total current iterations.

From Table (2), the high temperatures (i.e. $T=10000000000$) lead the SA to accept every worse solution. Such as, when the $\gamma$ is equal 150, 50 or 5. Namely, high temperature makes the SA to waste more computational time by accepting every worse bad solution. In this study, we select the initial temperature value based on the average deviation of the penalty value $\gamma$ (see Eq.1). However, according to the preliminary experiment, we suggest several initial temperatures ranges to select the suitable one among them according to the average deviation of the penalties value that will be calculated for several iterations during the SA process. The initial temperatures range in Table 3.

<table>
<thead>
<tr>
<th>$\gamma$</th>
<th>$T_0$</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 and more</td>
<td>Random (1000, 2000)</td>
</tr>
<tr>
<td>5 to 49</td>
<td>Random (500, 1000)</td>
</tr>
<tr>
<td>1 to 4</td>
<td>Random (100, 500)</td>
</tr>
</tbody>
</table>
Note: $\gamma$ is the deviation average of the penalties

<table>
<thead>
<tr>
<th>Technology</th>
<th>First Initial Temperature</th>
<th>Average(\gamma)</th>
<th>Comments</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dynamic</td>
<td>Dynamic</td>
<td>$</td>
<td>\gamma</td>
<td>$</td>
</tr>
<tr>
<td>SA with Estimated Temperature [16]</td>
<td>Infinite</td>
<td>$</td>
<td>\gamma</td>
<td>$</td>
</tr>
<tr>
<td>Solving the Course Scheduling problem Using SA [3]</td>
<td>10000</td>
<td>$</td>
<td>\gamma</td>
<td>$</td>
</tr>
</tbody>
</table>

Note $\bar{|\gamma|}$ means that the deviation average for (positive and negative values).

From Table 4, we found that the other researchers’ techniques have some limitations such as initializing high initial temperature values even when they used a small neighbors size, also the temperature becomes uncontrolled SA parameter anymore when it is estimated for each iteration, which means that the problem of setting the initial temperature value is still exists. Therefore, the dynamic Initial Temperature overcomes these problems by choosing the initial temperature value properly. Where the initial temperature $T_0$ is selected from the Table 2 suggested according to preliminary experiment from Table 1.

2.2 Adaptive Cooling Schedule (ACS)

The performance of SA depends heavily on temperature cooling schedule [5], where the good cooling schedule is substantial feature to get the optimal solution and to reduce the consuming time [17].

In this research, two cooling schedule proposed in the literature review are investigated in order to identify the good and suitable one as follow:

1. The first one is called Static or Geometric cooling schedules, which proposed by Kirkpatrick et al. [11] as the following equation:

   \[ T_{k+1} = \alpha T_k \]  
   
   where $(0 < \alpha < 1)$ denotes the cooling rate or factor; $T_k$ is the current initial temperature.
2. The second one is called Adaptive cooling schedules presented by Lewis et al. [12] where the temperature is decremented to be adjusted during the process of the algorithm according to the objective function values which is used to decide amount of temperature decreasing as following equations:

\[
\lambda_0 = 1 - \beta
\]

\[
\lambda_{i+1} = \lambda_i + \frac{\beta + \beta}{M}
\]

\[
T_{i+1} = T_i - \lambda_{i+1}\left(\frac{T_0}{M}\right)
\]

Where \(\beta\) represents a parameter to determine the value of \(\lambda\) which influence the concavity amount that presented in cooling schedule and \(M\) represents the iteration number used to decrease the initial temperature \(T_0\) to a zero value or close to it. However, this work investigates both geometric and adaptive cooling schedules, which were presented by Kirkpatrick et al. [11] and Lewis et al. [12] In order to identify the suitable cooling schedule for our study.

### 2.3 Neighborhood structure

Lü and Hao [13] claimed that neighbourhood structure is one of the most important features of the local search algorithms, where the effective and good neighbourhoods structure influence the SA performance positively [14]. Furthermore, Fleischer and Jacobso [9] stated that to improve the solution quality and the SA performance we must chose a good neighbourhood structure and size. Also, [7] mentioned that:

A smooth topology with shallow local minima which imposed by a neighbourhood structure is preferred, rather than a bumpy topology with many deeply local minima

Several researchers claimed that SA performance is better when the neighbourhood structure size is relatively small [6]. Meanwhile, other researchers demonstrated that SA performance improves when a large neighbourhood structure is used [15].

In this work, we use three neighbourhood structures in order to improve the solution quality, two neighbourhood structures are common (simple move as NS1 and simple swap as NS2), and a new proposed neighbourhood structure NS₃ which is presented by Tarawneh et al. [20].

**NS₃**: Generally, the selection mechanism in neighbourhood structure happens randomly (e.g.: - select two lectures randomly which belong to two different
rooms and slots). Thus, the search may need more time to reach a good solution. Tarawneh et al. [20] presented a new neighbourhood structure to reduce the random selection. \( NS_3 \) Neighbourhood structure mechanism calculates the total soft constraints penalties for each timeslot and sums them up for the whole week (Table 4 as example). Then, the lectures with the highest penalty swap with other lectures randomly.

Table (4) shows that the presented neighbourhood structure mechanism selects the timeslot with highest penalties (i.e. timeslot = three), and randomly select another timeslot. Then it interchanges both of them with any conflict. Therefore, the soft constraints violation weight (penalties) is calculated for each timeslot every time when a new neighbourhood structure is applied. However, this neighbourhood is used to minimize the random selection in order to avoid the disconnected neighbourhood structure that leads the search to escape from local optimum solution. it works as a flipping procedure for the SA search space.

| Table 5: The soft constraints penalties for each time period and slot |
|--------------------------|----------|----------|----------|----------|----------|--------|
|                         | Room1    | Room2    | Room3    | Penalties |
| Days (1-5)              | Days (1-5) | Days (1-5) | Days (1-5) |          |
| 0 5 6 30 0 0           | 100 1 1 4 2 | 10 20 0 1 160 | 339       |
| 6 16 10 0 0           | 2 4 2 30 100| 1 0 30 1 3 | 243      |
| 10 0 80 10 0         | 1 20 1 1 1 | 3 0 0 0 0 | 127      |
| 10 1 4 5 2           | 0 10 0 10 20| 6 4 150 1 10 | 233     |
| 1 2 8 1 10          | 20 30 15 0 1 | 2 2 6 0 0 | 165      |
| 2 3 6 1 8           | 120 30 40 0 4 | 0 0 9 0 1 | 224      |

Figure 5 shows the SA pseudo code, which contain D-IT, A-CS and the new neighbourhood structure.

Hint: for the parameters settings, we setup them for each part of our experiments separately.

3. Experiment and result

3.1 Experimental results of the Adaptive Cooling Schedule (ACS), D-IT with adaptive neighborhood structure

In the next experiment we compare the geometric cooling schedule (GCS) presented by Kirkpatrick et al. [11] with static initial temperature and standard neighbourhoods structure N1 and N2 against the adaptive cooling schedule (ACS) presented by Lewis et al. [12] with D-IT presented by Tarawneh et al. [20] and A-CS presented by Tarawneh and Ayob [19]. The parameters setting that we used in this experiment are presented in Table 6.

\( NS_1 \): Move one lecture period from the current period to another free position period.

\( NS_2 \): Randomly swaps two different lectures from different time slots and rooms.

\( NS_3 \): Adaptive neighborhood structure presented by Tarawneh et al. [20].
Table 6: SA parameters setting (standard and enhanced comparison)

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Standard SA(S-SA)</th>
<th>Enhanced SA(E-SA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial Temperature</td>
<td>1000000</td>
<td>D-TI</td>
</tr>
<tr>
<td>Neighborhoods structure</td>
<td>NS1 and NS2</td>
<td>NS3</td>
</tr>
<tr>
<td>Cooling rate</td>
<td>0.99</td>
<td>Adaptive</td>
</tr>
<tr>
<td>Termination condition</td>
<td>460 seconds</td>
<td>460 seconds</td>
</tr>
</tbody>
</table>

Table (6) illustrates the results of S-SA and E-SA. The first column indicates the instances, columns 2-5, 6-9 reports the best and worst solutions, Mean, and standard deviation over 31 runs for each instance and Column 10 reports the P-value.

Table 7. Computational results of the SA performance using standard components (S-SA) and Enhanced components (E-SA).

Table (7) shows that E-SA improves the SA performance compared to S-SA. SA permanence is statistically significant performance depending on enhanced simulated annealing components (in all instances except the small instances comp1 and comp11). Thus, we conclude that adaptive cooling schedule and
dynamic initial temperature with adaptive neighborhood structure will lead the SA to improve the solution quality compared to SA with standard and static components. *Note: Best results in italic bold.*

4. Conclusion

In this paper, we employed the standard SA components to solve the university course timetabling (UCT) problem using the benchmark dataset ITC2007-Track3. We investigated the SA components as follows: initial temperature, cooling schedule and neighborhood’s structure, by initializing the initial temperature dynamically and identifying a good cooling schedule to decrease the temperature properly. Moreover, we worked to improve the SA solution quality using adaptive neighborhood structure that swaps the selected time slot randomly with the timeslot that has the highest cost (penalty).

However, SA still could get stuck or trapped in local optimum, in the future work we suggest to hybridizing SA with other approaches or Artificial intelligence (AI) algorithms, in addition to test the solutions using a real world data set for further investigations.

References


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