Three-Stage Tabu Search for Solving Large-Scale Flow Shop Scheduling Problems

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Tabu search is a meta-heuristic approach designed skillfully for finding a suboptimal solution of combinatorial optimization problems. In this paper the tabu search with three stages is proposed for solving large-scale flow shop scheduling problems. In order to obtain a better suboptimal solution in a short computation time, three different candidate lists are used to determine the incumbent solution in the respective search stages. The candidate lists are constructed by restricting the moving of each job. Test problems with four kinds of job data are examined. Based on analyzing the relationship between the candidate list and the suboptimal solution for each job data, a common parameter is given to construct the candidate list during the search process. Comparison of the computation result is made with the genetic algorithm and the basic tabu search, from which it is shown that the proposed tabu search outperforms two others.

Keywords: Tabu search, Flow shop, Tardiness, Candidate list

1. Introduction

In manufacturing systems effective scheduling of jobs has become important for improving the efficiency of operation and decreasing the production cost. Tardiness criterion is of great importance because certain costs are incurred for a job not completed by its due date. These costs include: penalty clause in the contract, if there are any; loss of good will resulting in an increased probability of losing the customer for some or all future jobs, and a damaged reputation which will turn other customers away [1]. Flow shop problems viewed as sequence problems (called permutation flow shop problems) are concerned in this paper and the objective is to minimize the total tardiness $F \parallel \sum T_i$.

Tabu search (TS) was proposed by Glover [2], and aims at guiding the search beyond local optimum in the solution space of a given problem. In the real production field for large-scale and complicated scheduling problems we have to accept a suboptimal solution, because the computation time is limited. According to the paper written by Armentano [3] and Diaz [4], restricting the neighborhood of the current solution leads to reduction of computation time and improvement of the computational efficiency.

TS is one method of neighborhood search, and its performance depends on the neighborhood structure. Different neighborhoods have different properties, which may make them more or less suitable depending on the particular problem at hand. In our earlier paper [9], we proposed a tabu search with restricted neighborhood (TSRN) to solve flow shop problems. The knowledge obtained from the simulation of an adaptive behavior of fish schools has been used to define the neighborhood in TSRN. When the computation time is limited, the suitable neighborhood size has been given in TSRN for different kinds of job data.

We also proposed a tabu search with two stages (TSTS) [6] for solving the parallel machine problems $F \parallel \sum T_i$. In TSTS the heuristic method NEH is used to order the jobs assigned to each machine at the first stage, and the candidate list with small size is employed for reducing the search space at this stage. In order to rise the accuracy of the suboptimal solution obtained by the first stage, the intensive search with large candidate list is performed at the second stage. Because the search route is changed by changing the structure of candidate list, the risk of trapping in local minimum is reduced in TSTS.

In this paper a tabu search with three stages (TS3S) is proposed to minimize $F \parallel \sum T_i$. Because the solution contribution changes along with the search process, the different candidate lists are used in the three search stages for determining the incumbent solution. At the beginning there are many neighboring solutions better than the current solution, the random search is used and the candidate list with small size is employed to reduce the computation time. As the accuracy of solution rises, the number of better solutions included in the neighborhood set is reduced, and the large candidate list is used to improve a suboptimal solution.

The standard approach of intensification and diversification is able to improve the accuracy of the suboptimal solution. But the computational time is not acceptable
for large-scale problems. In TS3S new methods are used to determine the intensification and the diversification approach.

2. Flow Shop Problem

The permutation flow shop problem is formulated as follows. $n$ jobs $\{J_1, J_2, \ldots, J_n\}$ have to be processed in the same sequence on $m$ machines. The processing time of job $J_i$ on machine $M_j$ is given by $p_{ij}(i = 1, 2, \ldots, n, j = 1, 2, \ldots, m)$. These times are non-negative and some of them may be zero if the job is not processed on the machine. The due date of job $J_i$ is $d_i$.

For this problem the following assumptions are made:

- Every machine processes only one job at a time.
- Every job is processed on one machine at a time.
- Every job has to be processed at most once on machine $M_1, M_2, \ldots, M_m$ (in this order).
- The operations are not preemptive.

The objective function is to minimize the sum of the tardiness of each job. Let $s_{ij}$ and $c_{ij} (c_{ij} = s_{ij} + p_{ij})$ be the start time and the completion time of operation $O_{ij}$ of job $J_i$ processed on machine $M_j$, respectively.

The problem is to find a sequence $\pi = \{J_{i1}, \ldots, J_{in}\}$ so that the following objective function reaches a minimum value.

$$Z(\pi) = \sum_{i=1}^{n} \max(c_{im} - d_i, 0) \quad \cdots \quad (1)$$

3. Tabu Search

TS is a meta-heuristic approach which can be designed skillfully for finding a suboptimal solution of combinatorial optimization problems. Basically it consists of several components called the move, neighborhood, initial solution, attribute selection, tabu list, aspiration criterion, and stopping rules. TS starts from an initial solution. The move is a function which transforms a solution into another solution. The subset of moves applied to a given solution generates a collection of solutions called the neighborhood. The candidate list $C_L(\pi)$ is defined as the list of solutions evaluated in the neighborhood of the correct solution $\pi$. At each iteration a move leads to the best solution in $C_L(\pi)$ which may not be an improved solution. To avoid cycling and move generally to add robustness to the search, a recency-based short term memory called a tabu list $T = T_1, T_2, \ldots, T_n$ gives forbidden moves. A tabu list consists of attributes of the latest moves which are not allowed to be repeated during several iterations (the tabu tenure). But a forbidden move can be admitted if an aspiration criterion is satisfied.

The basic tabu search (BTS) is given as follows.

BTS

Step1: Set $t \leftarrow 1$ as an iteration number. Select the number $x$ from $[n/2, n]$ randomly. Set the initial tabu list $T_j = 0; j = 1, 2, \ldots, n$. $t_n \leftarrow 0$.

Step2: Generate the initial solution $\pi$. $\pi^* \leftarrow \pi$ and $\pi_{\text{best}} \leftarrow \pi$.

Step3: Construct the candidate list $C_L(\pi^*)$.

Step4: Obtain $Z(\pi_i) = \min\{Z(\pi_i), \pi_i \in C_L(\pi^*)\}$.

Step5: If $Z(\pi_i) < Z(\pi_{\text{best}})$ then $\pi_{\text{best}} \leftarrow \pi_i$, $t_n \leftarrow 0$, $T_1 \leftarrow x$, and go to Step 7. Otherwise find the best $\pi_i$ in $C_L(\pi^*)$ subject to $T_i = 0$. $t_n \leftarrow t_n + 1$.

Step6: If $Z(\pi_i) < Z(\pi_{\text{best}})$, then $T_i \leftarrow x - 1$. If $Z(\pi_i) \geq Z(\pi_{\text{best}})$, then $T_i \leftarrow x + 1$.

Step7: Renew the tabu list, $T_i \leftarrow T_i - 1$; $i = 1, 2, \ldots, l - 1, l + 1, \ldots, n$. $\pi^* \leftarrow \pi_i$.

Step8: If $t_n = n$, $\pi_{\text{best}}$ is adopted as the suboptimal solution and stop the computation. Otherwise $t \leftarrow t + 1$.

Step9: If the value of $t/20$ is an integer, update $x$ by an integer selected randomly from $n/2$ to $n$. Go to Step 3.

4. Components of the Proposed Tabu Search

The following components are adopted for the proposed TS.

Initial solution: At the initial sequence $\pi_{\text{init}}$, jobs are arranged in the non-decreasing order of due dates called EDD (earliest due date).

Move Among many types of moves considered in the literature for the flow shop problem, the following two types are used prominently:

Swapping Swap jobs placed at the $a$th and $b$th positions.

Insertion Remove a job placed at the $a$th position and insert it in the $b$th position.

In the proposed TS both types of moves are applied at the identical probability.

Neighborhood: For a given solution $\pi$, the solutions generated by swapping and inserting compose the swapping neighborhood $SN(\pi)$ and the insertion neighborhood $IN(\pi)$, respectively. For flow shop scheduling problems with $n$ jobs, the size of $SN(\pi)$ is $n(n - 1)/2$, and that of $IN(\pi)$ is $(n - 1)^2$. If the value of $n$ is large, we have heavy computation burden on checking the whole neighborhood to determine the incumbent solution.

Candidate List: The candidate list $C_L(\pi)$ is applied in order to reduce the effort for evaluating the neighborhood at each iteration of the search process. How to select the neighboring solutions from the neighborhood, and how many neighboring solutions should be selected to construct $C_L(\pi)$ are difficult problems for us. In the proposed TS3S three different $C_L(\pi)(i = 1, 2, 3)$ are used at the respective search stages.

Tabu list: The tabu list is defined as the set $T = T_1, T_2, \ldots, T_n$. $T_i$ is the tabu tenure of job $J_i$. $T_i$ can not be chosen for move until $T_i = 0$.

Tabu tenure: The tabu tenure is set dynamically. For every 20 iterations a new value for the tabu tenure is generated from a uniform distribution between $n/2$ and $n$, and at each iteration this value is adjusted. This adjustment is carried out according to the value of the selected solution (the incumbent solution) compared with the solution (the current solution) that generated the move. If the objective value of the incumbent solution is larger than that of the current solution, the tabu tenure of $J_i$ whose moving leads to the current solution translated to the incumbent solution,
is increased by one unit. Otherwise the tabu tenure of \(J_t\) is decreased by one unit. A similar procedure was proposed by Dell’Amico et al.\(^{(6)}\), and used by Armen- tano \(^{(3)}\).

**Aspiration criterion** A move can be released from the tabu condition if it has a better solution than the best solution found by the search until that moment.

**Stop rule** When the suboptimal solution is not improved after continuing \(n\) iterations, the search process is stopped at each stage.

5. Design of the Tabu Search with Three Stages

TS is one of neighborhood search algorithms. The starting point of the neighborhood search method is the choice of neighborhood structure; this determines which pairs of solutions are regarded as adjacent to each other. Consequently allowable moves are determined.

Small neighborhoods are preferable from the viewpoint of the speed at which the local search proceeds. If the neighborhood is too small, however, the final solution may be of poor quality. Neighborhood search techniques should be done in such a way that a move to an adjacent solution leads to a rapid update of the objective function value, rather than a complete recalculation based on the whole neighborhood.\(^{(9)}\)

In this paper our purpose is to propose a tabu search for solving large-scale flow shop problems. Instead of the whole neighborhood the candidate list is used for determining the incumbent solution. The construction of the candidate list is different for the respective stages.

5.1 The First Stage Process TS3S-I At the first stage, the search process begins from the initial solution which is obtained by EDD. Since at the beginning search stage we can easily find better neighboring solutions than the current solution, it is not necessary to recalculate all of the neighboring solutions to determine the incumbent solution. The candidate list with small size is used at this stage. The candidate list \(C_1^I(\pi)\) of the current solution \(\pi\) is constructed as follows:

**Construction of \(C_1^I(\pi)\)**

**Step1:** \(i \leftarrow 1\).

**Step2:** Job \(J_t\) is removed from the current position to the other arbitrary position in \(\pi\), or is swapped with an arbitrary job. The probabilities of insertion and swapping are 50%, respectively. The new solution is represented as \(\pi^t\). The move distance \(D_I(\pi, \pi^t)\) is given by:

\[
D_I(\pi, \pi^t) = |P_{\pi^t}^i - P_{\pi}^i| \quad \cdots \cdots \cdots \quad (2)
\]

where \(P_{\pi^t}^i\) and \(P_{\pi}^i\) are the position number of \(J_t\) in sequence \(\pi\) and \(\pi^t\), respectively.

**Step3:** If \(i = n\) stop; otherwise \(i \leftarrow i + 1\) and go to Step 2.

By this way there are \(n\) neighboring solutions included in \(C_1^I(\pi)\). At each iteration only \(n\) solutions are computed, and one solution \(\pi^t\) is selected as the incumbent solution for the next iteration. We define \(\delta_i(i = 1, 2, \ldots, n)\) as the number of moves whose distance is \(i\) during the search process. If at an iteration \(\pi\) translates to \(\pi^t\), and \(D_I(\pi, \pi^t)\) is the move distance of \(J_t\), then \(\delta_{D_I(\pi, \pi^t)} = \delta_{D_I(\pi, \pi^t)} + 1\).

5.2 The Second Stage Process TS3S-II In TS the best solution included in candidate list is selected as the incumbent solution. The EDD is used as the initial solution at the first stage. The solution generated with large \(D_I(\pi, \pi^t)\) is difficult to be selected as an incumbent solution, when the due dates of job data distribute in the wide time range. Since we focused on the random selection in the first stage, the distribution of \(\delta_i\) relates with the characteristic of the job data.

How can this information be used to predict which moves are likely to lead to improvements in the objective function value? The answer is that if we can restrict the move distance of job \(J_t\) in a suitable range, the evaluation of unnecessary non-improving moves is avoided and consequently search efficiency is improved. Therefore our purpose is to determine the suitable move range.

The parameter \(K\) is introduced to restrict the moving of job in the sequence, and the value of \(K\) is determined as the minimal value obtained from the following equation.

\[
\rho \leq \sum_{i=1}^{K} \delta_i / \sum_{i=1}^{n} \delta_i \quad \cdots \cdots \cdots \cdots \quad (3)
\]

We will give the value of \(\rho\) by analyzing the computation result later.

In our earlier paper\(^{(5)}\), the set \(C(\pi, J_t, I)\) is used to restrict the moving of each job when generating the neighboring solutions of the current solution. The definition of \(C(\pi, J_t, I)\) is given as follows:

\[
\sigma_{ik} = \begin{cases} 
\sigma_{k1} - \sigma_{k1} & \text{for } \sigma_{k1} \leq \sigma_{k1} \\
\sigma_{k1} - \sigma_{k1} & \text{for } \sigma_{k1} > \sigma_{k1}
\end{cases} \quad \cdots \cdots \cdots \quad (4)
\]

\[
C(\pi, J_t, I) = \{J_k \mid k \neq i, k \in \{k_1, k_2, \ldots, k_I\}\}
\]

\[
\sigma_{ik} = l = 1, 2, \ldots, I \text{ are the least } L \text{ values of } \sigma_{ik} \quad \cdots \cdots \cdots \quad (5)
\]

When \(\sigma_{ij} = \sigma_{ik}(j \neq k)\), select one job from jobs \(J_j\) and \(J_k\) randomly and include it in set \(C(\pi, J_t, I)\). Therefore the set \(C(\pi, J_t, I)\) consists of the \(I\) nearest jobs from job \(J_t\) in sequence \(\pi\). Fig. 1 shows an example of the Gantt chart for \((n, m) = (8, 3)\) and \(I = 5\). For \(J_t = J_7\) we have the set \(C(\pi, J_t, 5) = \{J_1, J_4, J_2, J_3, J_6\}\).

Depending on \(C(\pi, J_t, I)\) the candidate list \(C_2^I(\pi)\) is
constructed as follows:

Construction of $C^3_L(\pi)$

Step1: $i \leftarrow 1$.

Step2: Remove $J_i$ from the current position and place it at the position where an arbitrary job included in set $C(\pi, J_i, I)$ is placed, or swap $J_i$ with an arbitrary job included in set $C(\pi, J_i, I)$, if the probabilities of insertion and swapping are 50%, respectively.

Step3: If $i = n$ stop; otherwise $i \leftarrow i + 1$ and go to Step 2.

Because the $I$ nearest jobs are found forward and backward from $J_i$ in sequence $\pi$, when $I \geq 2 \times K$, the move whose distance is shorter than $K$ can be obtained by Step 2.

The candidate list $C^3_L(\pi)$ is composed with $n$ neighboring solutions. Because the moving of each job is restricted by the set $C(\pi, J_i, I)$, the search space in this stage is reduced.

In TS an unimproved solution can be selected as the incumbent solution in order to reduce the risk to trap in the local minimum. At the first stage the size of candidate list is small and the solutions included in candidate list are generated randomly. Therefore the solution in which a job is moved to an unreasonable position may be accepted at the first stage as an incumbent solution. At the second stage the jobs can be moved to only the $I$ nearest positions. If a job is moved out of this range at the first stage, it is difficult to be put back at the second stage. Based on the initial solution $\pi_1$ and the best solution $\pi^2_{best}$ obtained by the first stage, we generate the initial solution $\pi^2_1$ for the second stage by the following algorithm:

Algorithm A

Step1: $\pi^2_1 \leftarrow \pi^1_{best}$

Step2: Calculate the value of $D_i(\pi^1_1, \pi^2_i), (i = 1, 2, \ldots, n)$ from Eq. (2). $D_i(\pi^1_1, \pi^2_i)$ is the difference of the positions of $J_i$ in solutions $\pi^1_1$ and $\pi^2_i$.

Step3: $D_i \leftarrow \max\{D_i(\pi^1_1, \pi^2_i); i = 1, 2, \ldots, n\}$. If $D_i \leq K$, then stop.

Step4: Move job $J_i$ in the solution $\pi^2_1$ from its current position to the position where $J_i$ is placed in solution $\pi^1_1$. Save the new solution as $\pi^2_1$, and go to Step 2.

In TS the diversification strategy is used to drive the search into a new region of the solution space which has not been examined yet. The new feasible initial solution is needed to restart the search process. Two prominent methods are used to generate the new initial solution: multi-start approach and systematic approach. In the multi-start approach the solution is generated randomly. In this case it represents a schedule which has been examined earlier, and the process is repeated. The systematic approach is to generate the solution based on how often each job has occurred at each position so far. This frequency is stored in a matrix which has to be updated at each iteration.

In TS3S-II a new initial solution is generated by algorithm A based on the solutions $\pi^1_1$ and $\pi^2_{best}$. The candidate list in TS3S-II is different from that used in TS3S-I, and then the search is not repeated. Because only two solutions are recorded during the search process, the computation time is reduced.

5.3 The Third Stage Process TS3S-III

In this stage the purpose is to improve the best solution obtained by two former stages. In the case of using the small neighborhood, the final solution may be of poor quality. Therefore the number of solution included in the candidate list $C^3_L(\pi)$ is increased. Because the value of $I$ relates with the characteristic of job data, we also restrict the move of jobs when the candidate list $C^3_L(\pi)$ is constructed. The construction is as follows:

Construction of $C^3_L(\pi)$

Step1: $i \leftarrow 1$.

Step2: Remove $J_i$ from the current position and place it at the position where every job included in set $C(\pi, J_i, I)$ is placed, to generate set $\{\pi^{l+i}; l = 1, 2, \ldots, I\}$. $\pi^{l+i}$ is the $l$-th solution obtained by insertion of job $J_i$. Swap $J_i$ with every job included in set $C(\pi, J_i, I)$ to generate set $\{\pi^{l+i}; l = 1, 2, \ldots, I\}$. $\pi^{l+i}$ is the $l$-th solution obtained by swapping of job $J_i$.

Step3: If $i = n$ stop; otherwise $i \leftarrow i + 1$ and return to Step 2.

Then we have $C^3_L(\pi) = \{\pi^{l+i} \cup \pi^{l+i}; i = 1, 2, \ldots, n; l = 1, 2, \ldots, I\}$. In this stage an intensive search is performed in the reduced solution space, and the number of solutions included in candidate list increases with $I$.

5.4 Whole Algorithm of TS3S

Fig. 2 shows the flowchart of the whole algorithm for TS3S. In the TS3S the same procedure as BTS is used repeatedly at the three search stages, but candidate lists $C^i_L(\pi); i = 1, 2, 3$ are used at the respective stages. $\pi^1_1; \gamma = 1, 2, 3$ are the initial solutions for the respective stages. The solution $\pi^1_1$ generated by EDD is used as the initial solution of TS3S-I. The initial solution $\pi^2_2$ of TS3S-II is generated by algorithm A. The initial solution $\pi^3_3$ of TS3S-III is the better solution between $\pi^1_{best}$ and $\pi^2_{best}$. The suboptimal solution is obtained as the final solution $\pi^3_{best}$ of the third
stage.

6. Computational Experiment

An experiment was carried out to examine the performance of TS3S. The algorithm is coded in C and the experiments are performed on Compaq ProLiant ML330. CPU is Pentium 566MHZ.

6.1 Test Problem We generated randomly a number of instances with various due dates and processing times. The processing times of operations $p_{ij}$ are selected from discrete uniform distribution $[1, 100]$. The due dates of job data are uniformly distributed between $ML \times (1 - \tau - \frac{R}{2})$ and $ML \times (1 - \tau + \frac{R}{2})$, where $ML$ is a lower bound of the makespan, which is given by

$$ML = \max\{\frac{1}{\tau} \times (\sum_{i=1}^{m} p_{ij} + \min_{i=1}^{m} \sum_{j=1}^{m} p_{ij}) + \min_{i=1}^{m} \sum_{j=1}^{m} p_{ij}\} \cdots (6)$$

$\tau$ and $R$ are the tardiness factor and the range of due date ($100$), respectively. Fig. 3 shows the following four cases of distribution of due date with different $\tau$ and $R$.

**Case 1:** low tardiness factor ($\tau = 0.2$) and small due date range ($R = 0.6$)

**Case 2:** low tardiness factor ($\tau = 0.2$) and wide due date range ($R = 1.2$)

**Case 3:** high tardiness factor ($\tau = 0.4$) and small due date range ($R = 0.6$)

**Case 4:** high tardiness factor ($\tau = 0.4$) and wide due date range ($R = 1.2$)

The release date of job $J_i$ is uniformly distributed in the range $[0, d_i - \sum_{j=1}^{m} p_{ij}]$.

6.2 Computation Results by TS3S Ten test instances $\omega_i (i = 1, 2, \ldots, 10)$ with $m = 3$, $n = 200$ are generated for each case. Then 40 test instances are treated by TS3S, where the value $\rho$ is changed from 65% to 95%. For each instance $\omega_i$ ten different runs are performed with various $\rho$ by changing random numbers. At each run the objective values $Z(\pi^1_{\text{best}})$, $Z(\pi^2_{\text{best}})$, and $Z(\pi^3_{\text{best}})$ are recorded.

For each case the three values of instance $\omega_i$ obtained by the $t$-th run are denoted as $Z(\pi^\gamma_{\text{best}})_{i \rho}(\gamma = 1, 2, 3; i = 1, 2, \ldots, 10; t = 1, 2, \ldots, 10)$ and $\rho = 65\%, 70\%, \ldots, 95\%)$). Then their averages for ten runs are given by

$$Z(\pi^\gamma_{\text{best}})_{i \rho} = \frac{1}{10} \sum_{i=1}^{10} Z(\pi^\gamma_{\text{best}})_{i \rho} \cdots (7)$$

The average relative improvement obtained by TS3S-II with respect to TS3S-I, and that obtained by TS3S-III with respect to TS-II are given respectively by

$$R_{21} = \frac{1}{10} \sum_{i=1}^{10} \frac{Z(\pi^1_{\text{best}})_{i \rho} - Z(\pi^2_{\text{best}})_{i \rho}}{Z(\pi^1_{\text{best}})_{i \rho}} \cdots (8)$$

$$R_{32} = \frac{1}{10} \sum_{i=1}^{10} \frac{Z(\pi^2_{\text{best}})_{i \rho} - Z(\pi^3_{\text{best}})_{i \rho}}{Z(\pi^2_{\text{best}})_{i \rho}} \cdots (9)$$

The total improvement rate is given by

$$R_f = \frac{1}{10} \sum_{i=1}^{10} \frac{Z(\pi^1_{\text{best}})_{i \rho} - Z(\pi^3_{\text{best}})_{i \rho}}{Z(\pi^1_{\text{best}})_{i \rho}} \cdots (10)$$

The values $R_{21}$, $R_{32}$ and $R_f$ depend on $\rho$, but its dependency is omitted in their expression for simplicity of notation.

Tables 1 and 2 show the values of $R_{21}$ and $R_{32}$ for various $\rho$. From the two tables the approaches of TS3S-II and TS3S-III are not effective for the instances with the low tardiness factor (Cases 1 and 2). For Case 3 a little improvement ($R_{21}$) is obtained by TS3S-II as shown in Table 1. For these three cases the value of $\rho$ does not affect the values of $R_{21}$ and $R_{32}$. On the other hand, Case 4 corresponds to the problems with high tardiness and wide due date range. Consequently it is considered that Case 4 treats difficult problems. For such case the significant improvements of $R_{21}$ and $R_{32}$ are obtained, and the approaches of TS3S-II and TS3S-III are effective.

The relationship between $R_f$ and $\rho$ for Case 4 is shown in Fig. 4. When $\rho < 80\%$ the value of $R_f$ is improved as $\rho$ increases, but when $\rho > 80\%$ this trend is saturated. Table 3 shows the variation of the average CPU time taken by TS3S with $\rho$. Because the number of solutions included in the candidate list is increased with $\rho$, the cost of CPU time also increases with $\rho$. Based on Fig. 4 and Table 3, and also because $\rho$ does not affect the
results in Cases 1-3, we recommend $\rho = 80\%$ as the suitable value of an important parameter in TS3S.

Fig. 5 shows the convergence curve for one run of an instance in Case 4 with $\rho = 80\%$. In the first stage and the second stage, the considerable convergence speed is obtained though the size of the candidate list is small.

After continuing $n (= 200)$ iterations with no improvement in the objective value in TS3S-I, the search of TS3S-I attains a local optimum. At this time, the diversification is performed, which is to restart the search from a new initial solution. By using the new candidate list and the new initial solution in TS3S-II, the search escapes from the local optimum and converges on a solution better than the suboptimal solution obtained by TS3S-I. The suboptimal solution obtained by TS3S-II is improved by intensive search in TS3S-III.

6.3 Comparison with Other Algorithms

BTS proposed by Armentano and the genetic algorithm with search space reductions (RCGA) proposed by Zhao et al. are employed to treat the instances in the four cases. The same procedure as presented in Section 3 is used by BTS. Instead of the candidate list $C_L(\pi)$, the iterative neighborhood $I_N(\pi)$ is employed to determine the incumbent solution for the next iteration in BTS. It is shown in Zhao’s paper that RCGA outperforms the standard genetic algorithm for solving large-scale flow shop problems. Then we adopt RCGA as an example of genetic algorithms. The population size of RCGA is set to 500 and the algorithm stops after 5000 generations. Crossover rate $P_C$ is 1.0 and mutation rate $P_m$ is 0.05.

We test ten instances for each case. Each instance is treated ten times by BTS and RCGA with changing random numbers. Table 4 shows the average of the objective value obtained by each method for each case. For the instances in Cases 1 and 2, the difference of suboptimal solution among BTS, RCGA and TS3S is small. For such cases not so bad solutions can be also obtained by heuristic methods such as EDD or ERT (earliest release time). For Case 3 TS3S outperforms BTS and RCGA a little, and for Case 4 significant improvement is obtained by TS3S as compared with BTS and RCGA. For Case 3 and Case 4, the results obtained by heuristic methods are inferior.

Let $\bar{Z}_i^B$ and $\bar{Z}_i^G$ ($i = 1, 2, \ldots, 10$) be the average objective values obtained by BTS and RCGA for each instance, respectively. The average improvement rate of RCGA for BTS and that of TS3S with $\rho = 80\%$ for BTS are given, respectively, by

$$ R_{BTS/RCGA} = 10 \sum_{i=1}^{10} \frac{Z_i^B - Z_i^G}{Z_i^B} \cdots \cdOTS
Armentano also proposed a neighborhood restriction strategy in TS, by which 40% computation time is reduced for BTS, but the suboptimal solution is not better than that obtained by BTS. On the other hand, Table 5 shows that, as compared with BTS, TS3S attains 25~68% reduction in CPU time and almost same accuracy for Case 1 to 3. In addition for Case 4 TS3S outperforms BTS significantly from the viewpoints of CPU time and accuracy.

For further investigation, each of ten instances in Case 4, called Data 1 to Data 10, is treated 100 times by BTS, RCGA and TS3S with changing random numbers, and the result of statistics for each instance is obtained. The level of significance $\alpha$ is set to 0.05. In order to apply the 95% statistical test to these data, the following statistical values are defined for the differences in the suboptimal objective value:

$$Z_{BTS/TS3S} = \frac{\bar{Z}_B - \bar{Z}(\pi_{\text{best}})}{\sqrt{\frac{S^2_B}{n} + \frac{S^2_{\bar{Z}}}{n}}} \quad \ldots \ldots \ldots (13)$$

$$Z_{RCGA/TS3S} = \frac{\bar{Z}_G - \bar{Z}(\pi_{\text{best}})}{\sqrt{\frac{S^2_G}{n} + \frac{S^2_{\bar{Z}}}{n}}} \quad \ldots \ldots \ldots (14)$$

Where $\bar{Z}_B$, $\bar{Z}_G$ and $\bar{Z}(\pi_{\text{best}})$ are, respectively, the average objective values obtained by BTS, RCGA and TS3S for each data. The values $S^2_B$, $S^2_G$ and $S^2_{\bar{Z}}$ are the sampling variances of $\bar{Z}_B$, $\bar{Z}_G$ and $\bar{Z}(\pi_{\text{best}})$, respectively. In the same way, the statistical values $T_{BTS/TS3S}$ and $T_{RCGA/TS3S}$ are defined for the differences in the CPU time.

Table 6 shows the result of the 95% statistical testing for ten data. All the values in this table are larger than 1.96, which means that significant differences in both the suboptimal objective value and the CPU time exist between TS3S and BTS(or RCGA) under the assumption of normal distribution because of large sampling size. This shows that TS3S outperforms other two methods.

7. Conclusion

In this paper a three-stage tabu search has been proposed for solving large-scale flow shop scheduling problems. Because in TS3S different candidate lists are used during the search process, the risk of trapping into local minimum is reduced. Consequently the quality of suboptimal solution is improved. Small size candidate lists are employed by TS3S-I and TS3S-II, and then the computation time is saved.

The recommendation of an important parameter is given as $\rho = 80\%$ after analyzing the computation result. Based on $\rho$ and the move distance recorded in TS3S-I, the parameter $I$ can be determined, from which the search space is reduced in TS3S-II and TS3S-III.

Based on the computational experiences for four cases of test problems, TS3S is proved to be effective, especially for the difficult problems (Case 4), as compared with the genetic algorithm and the basic tabu search.

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References


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