FlowSort parameters elicitation based on categorization examples

CoDE-SMG – Technical Report Series

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Abstract: In multi-criteria sorting methods, it is often difficult for decision makers to precisely define their preferences. It is even harder to express them into parameters values. The idea of this work is to automatically find the parameters of a sorting model using classification examples in the contexts of traditional sorting and interval sorting. Interval sorting, i.e. the possible assignment of alternatives into several successive categories, is defined in this paper. The sorting method we are working with is FlowSort, which is based on the PROMETHEE methodology. Starting with an evaluation table and known allocations, we propose a heuristic based on a genetic algorithm (GA) to identify the weights, indifference and preference thresholds but also profiles characterizing the categories. We illustrate both the performances of the algorithm and the quality of the solutions on three standard datasets in both cases.

Keywords: Preference Learning; PROMETHEE; FlowSort; Interval Sorting; Genetic Algorithm.

1 Introduction

Multi-criteria decision aid (MCDA) has been an active research field for more than 40 years. In this context, possible decisions are simultaneously evaluated on multiple conflicting criteria. For instance, in the common example of buying a new car, one typically tries to minimize the cost and consumption while maximizing performances, comfort, etc. Obviously no real car would be the best on all those criteria. Therefore, the notion of optimal solution is most of time replaced by the idea of compromise solution [1].

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Different approaches have been proposed to tackle this kind of problem, among which we can distinguish three main families [2]: multi-attribute utility theory (MAUT) [3], outranking [4], and interactive methods [5].

In this article, we decided to work in the context of outranking methods. Those are based on pairwise comparisons of the alternatives. Among the best-known we may cite ELECTRE [6] and PROMETHEE [7].

Usually, three main multi-criteria problems are considered: ranking, sorting and choosing [8]. In the ranking problematic, alternatives are ranked from the best to the worst ones on the basis of a complete or partial order. For instance, the world academic institutions’ ranking on the basis of different scientific indicators. The sorting problematic is dedicated to the assignment of alternatives into predefined categories. For instance, sorting countries into risk categories on the basis of economical, financial and political indicators. Finally, the choice problematic addresses situations where a subset of the best possible alternatives has to be determined, such as identifying the best candidate for a given position in a hiring process.

Let us point out that different methods have been developed to sort in a multi-criteria context. Among them, we may cite UTADIS [9], PROAFTN [10], ELECTRE TRI [11] and FlowSort [12].

The traditional approach used to deal with sorting problems is to directly ask the Decision Maker to give central, or limiting, profiles defining the categories, but also weights and preference parameters for each criterion. But obtaining such information is often a cognitive complex task. Most of the time, the Decision Maker does not know himself what are exactly his preferences. Furthermore, they are hard to correctly formalize into weights, thresholds and profiles.

Henceforth, we will limit ourselves to the sorting method developed within the PROMETHEE methodology: FlowSort. In this paper, we will consider the problem inside out. Based on an existing categorization of alternatives, we will try to find the parameters of the sorting model that best replicate this information. This will be formalized as an optimization problem that will be solved using a genetic algorithm.

Let us point out that similar works have been proposed in the context of ELECTRE TRI [13], [14] and [15]. We may also quote MR-Sort from Sobrie [16] which aims to find the parameters of a simplified version of ELECTRE TRI. More recently, another contribution from Zheng considers the problem of learning the weights of the criteria [17]. To the best of our knowledge, no article has already addressed this question for FlowSort.

In this contribution we will consider the elicitation of FlowSort parameters in two cases: sorting and interval sorting. This will be detailed in section 2.

In section 3, we introduce PROMETHEE and FlowSort. In the third section, we describe the genetic algorithm we will use to solve the optimization problem. Then, in the section 5, we apply the algorithm on different standard datasets and compare the results with other methods. Finally, we investigate the performances of the algorithm with respect to the parameters tuning.

2 Sorting and interval sorting

In a traditional sorting problem, we have a set $C = \{c_1, c_2, ..., c_k\}$ of $k$ ordered categories. We assume that $c_i \succ c_{i+1}$: $c_i$ is preferred to $c_{i+1}$. Therefore $C_1$ is the best category and $C_k$ is the worst one. Each alternative $a$ is assumed to belong to a single category. For instance, in
the case of country risk, credit rating agencies evaluate the sovereign debt of each country. This problem is an ordered sorting problem because each debt is associated to a single category, and the categories are ordered (ex: AAA is better than AA).

Now, we consider an extension of the ordered sorting to partial information. Henceforth, we will call it "interval sorting". The idea of the interval sorting is to use partial information. For instance, it could be stated that a credit does not belong to categories $C_1$ and $C_4$, but there is an uncertainty between $C_2$ and $C_3$. So the categorization would give an information such that it is both $C_2$ an $C_3$ (without being able to further refine the assignment).

In the context of interval sorting, the categorization is an interval with an upper and lower category. Because the categories are ordered, the alternative also obviously belongs to the categories in between.

In the next section, we expose how the sorting will be computed using FlowSort, in both cases: standard and interval sorting.

### 3 Promethee and FlowSort

In this section, we briefly present PROMETHEE II and FlowSort. We refer the interested reader to [18] for a detailed description of PROMETHEE and to [19] for FlowSort.

Let $A = \{a_1, a_2, ..., a_n\}$ be a set of $n$ alternatives and let $F = \{f_1, f_2, ..., f_q\}$ be a family of $q$ criteria. The evaluation of alternative $a_i$ for criterion $l$ will be denoted by a real value $f_l(a_i)$.

For each pair of alternatives, let’s compute $d_l(a_i, a_j)$, the difference of $a_i$ over $a_j$ on criterion $l$.

$$d_l(a_i, a_j) = f_l(a_i) - f_l(a_j) \quad (1)$$

A preference function, denoted $P_l$, is associated to each criterion $l$. This function transforms the difference of alternatives’ evaluations $d_l(a_i, a_j)$ into a preference degree of the first alternative over the second one for criterion $l$. Without loss of generality, we consider that criteria have to be maximized. $P_l$ is defined as follows:

$$P_l : \mathbb{R} \rightarrow [0, 1] : x \rightarrow P_l(x) \quad (2)$$

such that:

- $\forall x \in \mathbb{R}^- : P_l(x) = 0$,
- $\forall x, y \in \mathbb{R}_0^+ : x \leq y \implies P_l(x) \leq P_l(y)$

There are different kinds of preference functions. Henceforth, we consider only the linear one (see figure 1) which is characterized by two parameters: an indifference and a preference threshold: $q_l, p_l$.

$$\pi_l(a_i, a_j) = P_l[d_l(a_i, a_j)] = \begin{cases} 0 & \text{if } d_l(a_i, a_j) \leq q_l \\ \frac{d_l(a_i, a_j) - q_l}{p_l - q_l} & \text{if } q_l < d_l(a_i, a_j) \leq p_l \\ 1 & \text{if } p_l < d_l(a_i, a_j) \end{cases} \quad (3)$$
Once $\pi_l(a_i, a_j)$ has been computed for all pairs of alternatives, we may define the aggregated preference degree of alternative $a_i$ over $a_j$ using the weights $w_l$ associated to each criterion $l$. Weights are assumed to be positive and normalized.

$$\pi(a_i, a_j) = \sum_{l=1}^{q} w_l \pi_l(a_i, a_j)$$

(4)

The last step consists in calculating the positive flow score denoted $\phi^+_A(a_i)$ and the negative flow score denoted $\phi^-_A(a_i)$ as follows:

$$\phi^+_A(a_i) = \frac{1}{n - 1} \sum_{x \in A} \pi(a_i, x)$$

(5)

$$\phi^-_A(a_i) = \frac{1}{n - 1} \sum_{x \in A} \pi(x, a_i)$$

(6)

We define the net flow score of $a_i$ as the difference between the positive flow and negative flows of $a_i$:

$$\phi_A(a_i) = \phi^+_A(a_i) - \phi^-_A(a_i)$$

(7)

The PROMETHEE I ranking is obtained as the intersection of the rankings induced by $\phi^+$ and $\phi^-$. For an interpretation of the net flow scores, the interested reader is referred to [20]. Finally, a complete order, called PROMETHEE II, can be derived from the order induced by $\phi$.

Based on PROMETHEE, FlowSort has been developed to address sorting problems [19]. Let $C = \{c_1, c_2, ..., c_k\}$ be a set of $k$ ordered categories. We assume that $c_i \succ c_{i+1}$: $c_i$ is preferred to $c_{i+1}$. Therefore $C_1$ is the best category and $C_k$ is the worst one.

Categories are assumed to be represented by limit or central profiles. On the one hand, the idea of the limiting profiles is to define couples of values for each criterion, defining the lower and upper bounds of the considered category. Let us note that the profile defining the upper bound of category $c_i$ is the same as the one defining the lower bound of category $c_{i+1}$.

On the other hand, central profiles are defined using a single value for each criterion. This
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represents a kind of mean profile of the category. A common property is that the profiles of each category must dominate the profiles of the ones they are preferred to. In this work, we have chosen to work with central profiles.

Let’s define \( R = \{r_1, r_2, \ldots, r_k\} \), the set of central profiles representing the \( k \) categories. To identify the category of an alternative \( a_i \), we define the subset \( R_i = R \cup \{a_i\} \). Then, for each element \( x \) in the subset \( R_i \), we compute its net flow score \( \phi_{R_i}(x) \).

As in the nearest neighbor procedure, the category of alternative \( a_i \) is the one such that the profile has its net flow score the closest to the net flow score of \( a_i \). More formally:

\[
I^*(a_i) = \arg\min_{l=1,2,\ldots,k} |\phi_{R_i}(a_i) - \phi_{R_i}(r_l)|
\]

Let us note that \((3 + k) \cdot q\) parameters have to be provided in order to instantiate FlowSort:

• \( k \cdot q \) values for the central profiles;
• \( 3 \cdot q \) values for the weights, indifference and preference thresholds.

In the case of the interval sorting problem, an alternative can be sorted in multiple consecutive categories. In this case, we use an extension of PROMETHEE I instead of PROMETHEE II. The upper and lower categories are determined using the positive and negative flow scores. As in the regular FlowSort method, the category of \( a_i \) is determined as the category of the profile having its positive, resp. negative, flow score the closest to the one of the alternative \( a_i \).[19]

\[
I^+_i(a_i) = \arg\min_{l=1,2,\ldots,k} |\phi_{R_i}^+(a_i) - \phi_{R_i}^+(r_l)|
\]

\[
I^-_i(a_i) = \arg\min_{l=1,2,\ldots,k} |\phi_{R_i}^-(a_i) - \phi_{R_i}^-(r_l)|
\]

If both values are equal, the categorization is precise. Otherwise, these values define the range of the categories.

4 Algorithms

As a first approach, we propose to use a genetic algorithm (GA) to address the elicitation problem. The use of a metaheuristic is motivated by the non linear nature of preference functions. In this work, we have implemented the algorithms using jMetal; a Java Framework that contains already a lot of well-known optimization algorithms [21, 22]. From our point of view, the added value of jMetal is the design of the optimization problem. It allows to be very flexible with the algorithm. It is indeed easy to compare different mutation and crossover operators, etc. The pseudo code of the algorithms described in this section is available in the appendices of this article.

Additionally, we used iRace\(^3\) which is a tool that implements the iterated racing procedure, in order to fine tune the parameters of the GA [23] [24] in the case of standard sorting. For the interval sorting, we used another method which changes the value of each parameter with respect to the optimization process.

The algorithm is quite the same for both cases (standard sorting and interval sorting), the only differences are in the evaluation method of the distance and correctness of a solution.

Each solution contains, for each criterion \( l \):
• a weight $w_l$;
• an indifference threshold $q_l$;
• a preference threshold $p_l$;
• the $k$ values $r_{hl}$, $h \in \{1,\ldots,k\}$ defining the categories profile for each criterion.

For convenience, we assume that all the evaluations of the alternatives have been normalized on a scale $[0,1]$.

The structure of our genetic algorithm is quite common (see algorithm 1). We begin by generating an initial population. We set a limit on the number of evaluations denoted $\text{max} \_ \text{eval}$. This will be used as a stopping criterion. Obviously, the algorithm also stops when a perfect solution is found. A new population is created by selecting parents (with a selection operator based on quality) and creating offspring with particular crossover and mutation operators. Let us note that, for each generation, the two best solutions at this stage of the optimization process are kept in the new population.

A major step is to define an indicator that will quantify the quality of a given solution. Considering the case of standard sorting, we have chosen to use the $L_1$ distance between the categorization given by FlowSort with the actual parametrization and the categorization given as input:

$$f(s) = \sum_{a \in A} |l^*_s(a) - l(a)|$$

(11)

where $l(a)$ represents the known categorization of the alternative $a$ and $l^*_s(a)$ the categorization using the parameters of solution $s$. The distance is used to induce a higher penalty if there is a big difference between the prediction and the real category.

A distinctive feature in the case of interval sorting is that we have to deal with two pieces of information: the upper and lower categories for each alternative. Let us note $c^+(a_i)$ the upper category and $c^-(a_i)$ the lower one. We use here the $L_1$ distance between the upper and lower category given as input $c_r$ and the one given by the current parametrization of FlowSort $c_f$. Intuitively, the penalty associated to $s$ is defined as follows:

$$f(s) = \sum_{a \in A} (|c^+_f(a) - c^+_r(a)| + |c^-_f(a) - c^-_r(a)|)$$

(12)

For the sake of simplicity, we denote $c^+_f(a) = c^+_f(a,s)$. The same applies to $c_r$ and $c^-$. We define also the correctness of a solution $s$. The correctness defines how good a solution is with respect to the real categorization.

For the standard sorting the correctness is defined as the percentage of categories well categorized.

For the interval sorting, the correctness of a single alternative is defined as the number of categories correctly predicted divided by the total range covered by the predicted and the real categories. The correctness of a solution $s$ is defined as the sum of the correctness of all the alternatives:

$$\sum_{a \in A} \frac{\max(\min(c^+_f(a), c^+_r(a)) - \max(c^-_f(a), c^-_r(a)), -1) + 1}{\max(c^+_f(a), c^+_r(a)) - \min(c^-_f(a), c^-_r(a)) + 1}$$

(13)

In order to drive the search process to interesting solutions we define six indicators for each category. Let us denote them $U_l$, $W_l$ and $O_l$ for each category $l$: 
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- \( W_l \) represents the percentage of alternatives from the category \( l \) that are sorted in the same category using FlowSort;
- \( U_l \) represents the percentage of alternatives from the category \( l \) that are sorted in a lower category using FlowSort;
- \( O_l \) represents the percentage of alternatives from category \( l \) that are sorted in an upper category using FlowSort.

In a similar way, we define \( U^*_l \), \( W^*_l \) and \( O^*_l \) but this time using the categorization using FlowSort as the benchmark:

- \( W^*_l \) will represent the percentage of alternatives sorted in \( l \) that are really categorized in \( l \);
- \( U^*_l \) represents the percentage of alternatives sorted in \( l \) that are undercategorized, meaning their real category is higher than \( l \);
- \( O^*_l \) represents the percentage of alternatives sorted in \( l \) that are overcategorized, meaning their real category is lower than \( l \).

The initialization of the algorithm, see algorithm 2, is set up as follow:

- weights \( w_l \) are generated by using uniform random values between 0 and 1;
- indifference thresholds \( q_l \) are generated by using random values between 0 and 0.5;
- preference thresholds \( p_l \) are generated by using random values between \( q_l \) and 1;
- for the categories’ profiles, the range is divided in \( \text{nb
d_categories} \) equal parts. The value of a profile \( l \) on a criterion \( j \), denoted \( r_{lj} \), is chosen randomly between the value of the previous profile, or 0 if there is no, and the upper bound of the range.

The implementation follows a traditional GA scheme. Parents are selected using a binary roulette wheel selection based on the quality indicator. Depending on a crossover probability, two children are built based on linear combinations (each component depending on gene crossover probability).

The mutation step is based on the six indicators defined before. First, if the assignment correctness of a category is high, the mutation probability of its profile will be low (and reversely). Secondly, in case of unbalanced assignment mistakes, the profiles will be mutated:

- in ascending order if alternatives are too much overcategorized;
- in descending order if alternatives are too much undercategorized.

When a new solution is built, a correction is applied to ensure that profiles respect the dominance condition.

We refer the interested reader to the appendices for a complete presentation of the algorithms.
Table 1  Datasets properties for standard sorting.

<table>
<thead>
<tr>
<th>dataset</th>
<th>#inst.</th>
<th>#crit.</th>
<th>#cat.</th>
<th>% imprecise cat.</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU</td>
<td>209</td>
<td>6</td>
<td>4</td>
<td>40.67</td>
</tr>
<tr>
<td>BC</td>
<td>278</td>
<td>7</td>
<td>2</td>
<td>23.02</td>
</tr>
<tr>
<td>CEV</td>
<td>1728</td>
<td>6</td>
<td>4</td>
<td>16.43</td>
</tr>
</tbody>
</table>

Table 2  Datasets properties for interval sorting.

<table>
<thead>
<tr>
<th>dataset</th>
<th>#inst.</th>
<th>#crit.</th>
<th>#cat.</th>
<th>% imprecise cat.</th>
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<td>278</td>
<td>7</td>
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</tr>
<tr>
<td>CEV</td>
<td>1728</td>
<td>6</td>
<td>4</td>
<td>16.43</td>
</tr>
</tbody>
</table>

5  Results

In order to investigate the performances of our algorithm we have chosen to test it on three different benchmark datasets. This was only possible for standard sorting. Indeed, at the best of our knowledge, there has been no benchmark dataset for interval sorting yet. The datasets come from the website of Marburg University\(^4\). These originally come from the UCI repository\(^5\) and the WEKA machine learning toolbox\(^6\). The alternatives with missing values have been removed, ordinal values are transformed to numeric one, and the data have been normalized between 0 and 1. They are monotone learning datasets, each criteria has to be maximized. One can find the properties of the datasets in table 1. We have chosen the following three datasets: CPU, BC and CEV. Let us point out that these datasets have also been used by Sobrie [16] in the context of sorting but using a simplified version of ELECTRE TRI.

In the case of interval sorting, we decided to use the same 3 datasets but generating an interval categorization by using a random instantiation of FlowSort. Therefore we know an exact solution exists for the model’s parameters, which gives some kind of bias. The properties of the datasets are in table 2. The percentage of imprecise categorization represents the amount of alternatives that are sorted in an interval of categories instead of a precise category.

5.1  Sorting

In the case of standard sorting, we have used iRace\(^7\) [23] to fine tune of the parameters of the algorithm. In what follows, we will quantify the improvement gained with the use of iRace. The 5 parameters that have been optimized are: population size, mutation probability, gene mutation probability, crossover probability, gene crossover probability.

We have tested the algorithms as follow: each dataset has been divided in a learning set and a test set. Different size of learning set have been considered. Alternatives in the learning set have been randomly selected, but we have forced the algorithm to at least randomly select one alternative from each category. For each learning and test set, the algorithm has been executed on the learning set to elicit the parameters. Then the parameters found have been evaluated on the test set. This operation has been executed 16 times for each learning set. For robustness’ sake, the whole operation has been executed 10 times for each learning set size.
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Table 3 Results of the algorithm - correctness (standard sorting)

<table>
<thead>
<tr>
<th>learning set's size</th>
<th>dataset</th>
<th>r_init</th>
<th>p_init</th>
<th>p_rand_par</th>
<th>p_iRace</th>
<th>r_iRace</th>
</tr>
</thead>
<tbody>
<tr>
<td>5%</td>
<td>CPU</td>
<td>0.426 ± 0.076</td>
<td>0.671 ± 0.074</td>
<td>0.791 ± 0.079</td>
<td>0.681 ± 0.074</td>
<td>0.712 ± 0.087</td>
</tr>
<tr>
<td></td>
<td>BC</td>
<td>0.699 ± 0.036</td>
<td>0.687 ± 0.064</td>
<td>0.646 ± 0.079</td>
<td>0.694 ± 0.042</td>
<td>0.699 ± 0.060</td>
</tr>
<tr>
<td></td>
<td>CEV</td>
<td>0.700 ± 0.003</td>
<td>0.697 ± 0.018</td>
<td>0.723 ± 0.079</td>
<td>0.736 ± 0.041</td>
<td>0.777 ± 0.025</td>
</tr>
<tr>
<td>20%</td>
<td>CPU</td>
<td>0.336 ± 0.080</td>
<td>0.718 ± 0.084</td>
<td>0.862 ± 0.042</td>
<td>0.862 ± 0.048</td>
<td>0.876 ± 0.042</td>
</tr>
<tr>
<td></td>
<td>BC</td>
<td>0.716 ± 0.025</td>
<td>0.711 ± 0.032</td>
<td>0.720 ± 0.031</td>
<td>0.716 ± 0.027</td>
<td>0.716 ± 0.027</td>
</tr>
<tr>
<td></td>
<td>CEV</td>
<td>0.700 ± 0.006</td>
<td>0.696 ± 0.016</td>
<td>0.701 ± 0.099</td>
<td>0.772 ± 0.027</td>
<td>0.805 ± 0.013</td>
</tr>
<tr>
<td>35%</td>
<td>CPU</td>
<td>0.346 ± 0.068</td>
<td>0.763 ± 0.059</td>
<td>0.877 ± 0.088</td>
<td>0.920 ± 0.030</td>
<td>0.916 ± 0.029</td>
</tr>
<tr>
<td></td>
<td>BC</td>
<td>0.721 ± 0.020</td>
<td>0.714 ± 0.027</td>
<td>0.731 ± 0.027</td>
<td>0.738 ± 0.031</td>
<td>0.722 ± 0.021</td>
</tr>
<tr>
<td></td>
<td>CEV</td>
<td>0.703 ± 0.009</td>
<td>0.696 ± 0.016</td>
<td>0.705 ± 0.097</td>
<td>0.780 ± 0.040</td>
<td>0.812 ± 0.016</td>
</tr>
<tr>
<td>50%</td>
<td>CPU</td>
<td>0.350 ± 0.087</td>
<td>0.788 ± 0.068</td>
<td>0.897 ± 0.043</td>
<td>0.941 ± 0.020</td>
<td>0.943 ± 0.027</td>
</tr>
<tr>
<td></td>
<td>BC</td>
<td>0.727 ± 0.028</td>
<td>0.700 ± 0.029</td>
<td>0.723 ± 0.035</td>
<td>0.723 ± 0.033</td>
<td>0.736 ± 0.025</td>
</tr>
<tr>
<td></td>
<td>CEV</td>
<td>0.706 ± 0.011</td>
<td>0.700 ± 0.017</td>
<td>0.708 ± 0.094</td>
<td>0.781 ± 0.036</td>
<td>0.812 ± 0.015</td>
</tr>
</tbody>
</table>

Figure 2 Correctness with respect to the learning set size. Using the algorithm with parameters optimized with iRace, 250000 evaluations per learning set. (standard sorting)

Figure 3 Testing scheme (standard sorting)

A summary of the main results of our testing is available in table 3. The number represents the mean correctness of the prediction rate for the test set and the standard
deviation on this value. Here is a description of the different algorithm configurations (see figure 3):

- r_init. is a test based on the random generation of initial solutions (algorithm 2), we have generated 10000 initial solutions and selected the best one. No optimization is performed at this stage;

- p_init. is a random generation based on the random selection of an alternative for each category, the values are then sorted regarding each criterion to respect the dominance between the profiles. It is a generation of solutions using the information of the learning set. We have also generated 10000 solutions. No optimization is performed at this stage;

- p_rand_par. the initial solutions are generated the same way as p_init., but for the optimization we have randomly instantiated the parameters of the GA. We have limited the number of evaluations to 250000;

- p_iRace the initialization algorithm is the same as p_init. In this case, we have optimized the parameters of the GA using iRace (with 6000 evaluations). We have limited the number of evaluations to 250000;

- r_iRace is exactly the same as p_iRace but with the initial generation of solutions being random, like in r_init.

The results show a good prediction degree for the dataset CPU. We can also note an improvement on the solution quality for CEV. The gain on the correctness is clear with the parameters optimized using iRace instead of a random instantiation. Due to the nature of the BC dataset, no improvement is observed in table 3. The average performance of the proposed algorithm is as good as the one proposed for other methods, like MR-Sort for ELECTRE TRI or LP UTADIS (see table 4) [25]. These statistics have to be considered cautiously since experiments have not been executed with the same settings. Nevertheless, they allow to show that we reach similar performances.

The execution time is rather small, for instance for the CPU dataset the algorithm runs in about 18 seconds on a Intel i7-2640m with 8go RAM, running under Windows 8.1 with an implementation of the algorithm in Java 8.

In figure 2, one can see a boxplot representing the evolution of the correctness of the prediction rate in function of the size of the learning set. It is interesting to observe that we already reach a good prediction rate with low size of the learning set.

<table>
<thead>
<tr>
<th>learning set's size</th>
<th>dataset</th>
<th>META MR-SORT</th>
<th>LP UTADIS</th>
<th>r_iRace</th>
</tr>
</thead>
<tbody>
<tr>
<td>20%</td>
<td>CPU</td>
<td>0.7443 ± 0.0509</td>
<td>0.8679 ± 0.0488</td>
<td>0.876 ± 0.043</td>
</tr>
<tr>
<td></td>
<td>BC</td>
<td>0.7196 ± 0.0302</td>
<td>0.7085 ± 0.0307</td>
<td>0.699 ± 0.060</td>
</tr>
<tr>
<td></td>
<td>CEV</td>
<td>0.7761 ± 0.0183</td>
<td>0.7719 ± 0.0153</td>
<td>0.777 ± 0.025</td>
</tr>
<tr>
<td>50%</td>
<td>CPU</td>
<td>0.8092 ± 0.0361</td>
<td>0.9340 ± 0.0266</td>
<td>0.943 ± 0.021</td>
</tr>
<tr>
<td></td>
<td>BC</td>
<td>0.7246 ± 0.0446</td>
<td>0.7146 ± 0.0246</td>
<td>0.736 ± 0.025</td>
</tr>
<tr>
<td></td>
<td>CEV</td>
<td>0.7833 ± 0.0180</td>
<td>0.7714 ± 0.0158</td>
<td>0.812 ± 0.015</td>
</tr>
</tbody>
</table>

Table 4: Results of MR-SORT and LP UTADIS from [25] wrt r_iRace- correctness
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Figure 4  Varying parameters for the GA

<table>
<thead>
<tr>
<th>parameter</th>
<th>$p_{min}$</th>
<th>$p_{max}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>mutation probability</td>
<td>0.1</td>
<td>0.9</td>
</tr>
<tr>
<td>gene mutation probability</td>
<td>0.25</td>
<td>0.99</td>
</tr>
<tr>
<td>crossover probability</td>
<td>0.1</td>
<td>0.9</td>
</tr>
<tr>
<td>gene crossover probability</td>
<td>0.25</td>
<td>0.99</td>
</tr>
</tbody>
</table>

Table 5  Values of $p_{min}$ and $p_{max}$ for each parameter.

5.2 Interval sorting

First of all, let us mention that the results related to interval sorting have been published in the proceedings of the workshop DA2PL 2014.

Now considering the case of interval sorting, the fine tuning of the parameters we made was quite different.

With a closer look on the GA, we can see that this algorithm has mainly two kinds of exploration: diversification with the mutation operator and intensification with the crossover operator. During the tests, we have observed that the algorithm should ideally enforce diversification after intensification and then go back to intensification, and so on. As a consequence, the idea we have applied is to force the parameters variation of the algorithm during the optimization process. There are still our 5 parameters: population size, mutation probability, gene mutation probability, crossover probability, gene crossover probability.

The population size has been fixed to 1600 solutions. This value has been set after a set of trial and errors, and seems to work well in the considered examples.

The 4 others parameters have values between 0 and 1. At each step of the optimization we change the values of those following a linear equation. When the value 0, or 1, is reached the coefficient is reversed. We paid attention to chose different coefficients, so that the period is different. This permits to have a lot of different combinations of intensification and combinations.

In table 5, we show the values we have chosen for $p_{min}$ and $p_{max}$ for each parameter.

With this method, we have seen a slight improvement of the results for the standard sorting. We were able to increase the correctness of the prediction applied to the learning set. Unfortunately, this did not really improve the prediction rate on the test set using the
parameters learned with the learning set. However, let us note that this procedure is more simple than iRace.

The testing procedure has been set as follows: each dataset has been divided in a learning set and a test set. Different sizes of learning set have been considered. Alternatives in the learning set have been randomly selected. Nevertheless, we forced the algorithm to select randomly at least one alternative from each category. For each learning and test set, the algorithm has been executed on the learning set to elicit the parameters. Then the values found have been evaluated on the test set. This operation has been executed 32 times for each learning set. For robustness’ sake, the whole operation has been executed 10 times for each value of the learning set size. The maximum number of evaluations has been set to 2500000, and the population size to 1600. Results are available in table 6.

The correctness represents the accuracy of the prediction in the test set, and the learning set correctness represents the accuracy of the model on the learning set. From a global point of view, we can note that the correctness values are rather good. As expected, the correctness is increasing with the learning set’s size. One can note that the learning set correctness is decreasing with the learning set size too. This is because it is much more complicated to have a perfect solution if the number of alternatives in the learning set increases. For the dataset CEV, the results show that the algorithm does not reach good level of learning set correctness (at least not as high as for the two other ones). The reason probably lies in the fact that this specific dataset is much bigger. We have noticed that the performances on the test set are better than on the learning set. Currently, we have no explanation for this effect.

On figure 5, we show a boxplot of the evolution of correctness with respect to the learning set size (for the CPU dataset). We can note that a good level of correctness is already reached for a learning size of 20%.

Due to the introduction of the varying parameters for the GA, we did not need to fine tune the parameters anymore. The value of 1600 for the population size has been determined by trial and errors. Let us stress that the new method increases the running time. Nevertheless, it remains rather small. For instance, for the CPU dataset, the algorithm runs in about 5 minutes on a Intel i7-2640m with 8GB RAM, under Windows 8.1 with an implementation of the algorithm in Java 8. One advantage we have remarked during these first experiments is that the algorithm seems to be less stuck in local optima.
Figure 5  Correctness of the test set wrt LS size - CPU dataset (interval sorting)

6 Conclusion

In this paper, we have proposed a first method for the parameters elicitation of FlowSort in the case of standard sorting and interval sorting. The method is based on categorization given by the decision maker. Our approach is based on a genetic algorithm. The first results obtained are encouraging in both cases.

For instance, in the case of standard sorting, the running time is below 30 seconds for the CPU dataset containing 209 alternatives, 6 criteria and 4 categories. The performances of the algorithm on this instance is quite good, and we reach a good prediction rate of around 94%. The fine tuning of the parameters has been realized with iRace. In the case of interval sorting, we also obtain good results (but with another method for the fine tuning of the parameters). The parameters variation we use was able to balance the need of intensification and diversification the GA needed to be less stuck in local optima. But it can be noted that for both methods, the algorithm lead to a good overall result, which give an indication that it is able to converge to good solutions in both cases.

In further research, we may investigate the possibility of developing an exact method by using a simplified version of FlowSort.

Another interesting question is to improve the performances of the algorithm for small learning sets. In this context, a lot of very different solutions can lead to an exact prediction on the learning set but not on the test set. The combination of different instantiations of FlowSort can be investigated in order to improve the robustness of the allocation.
From an algorithmic point of view, a detailed analysis of the heuristic is still to be done. More precisely, quantitative arguments have to be highlighted in order to confirm the added value of parameters variations.

The use of benchmark datasets, in the case of interval sorting, (that are not linked to a particular method like in this study) will certainly have an impact on the prediction quality. Nevertheless, the existence (or the creation) of such datasets is far from being obvious.

Another question behind the interval sorting is the meaning of such imprecision. Is an alternative categorized in an interval of categories because the information we have is not complete enough but the alternative belongs only to a single category. Or is the alternative somehow in between two categories and whatever the amount of information we have we could not decide whether it is one or the other, because it could belong to both.

Finally, a more general question about the representability of FlowSort has to be deepened, which allocations can, or not, be reproduced by using such a model? This is also related to the identification of possible inconsistencies in the datasets.

Acknowledgement

We thank Olivier Sobrie for the constructive discussion in the beginning of this work.

A Algorithms

```
population = generate_initial_solutions();
while max_eval not reached do
    new_population = select_2_best_solutions(population);
    while size(new_population) lesser than size(population) do
        parents = selection_operator(population);
        new_solutions = crossover_operator(parents);
        mutation_operator(new_solutions);
        evaluate(new_solutions);
        new_population.add(new_solutions);
    end
    population = new_population;
    if population contains perfect solution then
        break;
    end
end
```

Algorithm 1: Genetic Algorithm structure

References

new_sol = create_sol_structure();
for $j = 1$ to $nb\_criteria$ do
    new_sol.w[j] = rand(0, 1);
    new_sol.q[j] = rand(0, 0.5);
    new_sol.p[j] = rand(new_sol.q[j], 1);
    new_sol.profile[1][j] = rand(0, 1/nb\_categories);
    for $l = 2$ to $nb\_categories$ do
        new_sol.profile[l][j] = rand(new_sol.profile[l-1][j], l/nb\_categories);
    end
end

Algorithm 2: Initialization operator

**input** : the *solution* to mutate; mutation probability
rand_nbr = rand(0, 1);
if rand_nbr < mutation_probability then
    rand_nbr = rand(0, 1);
    if rand_nbr < 0.4 then
        profiles\_mutation\_operator(solution);
    else
        if rand_nbr < 0.7 then
            weights\_mutation\_operator(solution);
        else
            thresholds\_mutation\_operator(solution);
        end
    end
end

Algorithm 3: Mutation operator
Algorithm 4: Mutation operator for profiles

```
input : the solution to mutate; gene mutation probability (gmp); \( U_l, W_l, O_l, U_l^*, W_l^*, O_l^* \) for each profile \( l \)

For simplifications purposes, we consider \( \forall j \text{ solution.profile}[l][j] = 0 \) for \( l < 1 \) and \( \text{solution.profile}[l][j] = 1 \) for \( l > \text{nb_categories} \).

for \( l = 1 \) to \( \text{nb_categories} \) do
    if \( \text{rand}(0, 2) > (W_l + W_l^*) \) then
        if \( U_l/(U_l + O_l) > 0.8 \) and \( W_l < 0.6 \) then
            for \( j = 1 \) to \( \text{nb_criteria} \) do
                \( \text{solution.profile}[l][j] = (\text{solution.profile}[l][j] + \text{solution.profile}[l-1][j]) / 2; \)
            end
        else
            if \( O_l/(U_l + O_l) > 0.8 \) and \( W_l < 0.6 \) then
                for \( j = 1 \) to \( \text{nb_criteria} \) do
                    \( \text{solution.profile}[l][j] = (\text{solution.profile}[l][j] + \text{solution.profile}[l+1][j]) / 2; \)
                end
            else
                for \( j = 1 \) to \( \text{nb_criteria} \) do
                    if \( \text{rand}(0, 1) < \text{gmp} \) then
                        if \( \text{rand}(0, 1) < U_l + U_l^* / (U_l + O_l + U_l^* + O_l^*) \) then
                            \( \text{solution.profile}[l][j] = \text{solution.profile}[l][j] + (U_l + U_l^*)/2 * \) \( \text{rand}(-\text{solution.profile}[l-1][j] - \text{solution.profile}[l][j], 0); \)
                        else
                            \( \text{solution.profile}[l][j] = \text{solution.profile}[l][j] + (O_l + O_l^*)/2 * \) \( \text{rand}(0, \text{solution.profile}[l+1][j] - \text{solution.profile}[l][j]); \)
                        end
                    end
                end
            end
        end
    end
end
```

Algorithm 5: Mutation operator for weights

```
import : the solution to mutate; correctness of the solution; gene mutation probability (gmp)

for \( j = 1 \) to \( \text{nb_criteria} \) do
    if \( \text{rand}(0, 1) < \text{gmp} \) then
        \( \text{solution.w}[j] = \text{solution.w}[j] + (1 - \text{correctness}) * \text{rand}(-\text{solution.w}[j], 1 - \text{solution.w}[j]); \)
    end
end
```
input : the solution to mutate; correctness of the solution; gene mutation probability ($gmp$)

for $j = 1$ to $nb\_criteria$ do
    if $\text{rand}(0, 1) < gmp$ then
        $\text{solution.q}[j] = \text{solution.q}[j] + (1 - \text{correctness}) \times \text{rand(}-\text{solution.q}[j], $\text{solution.p}[j] - \text{solution.q}[j])$;
    end
    if $\text{rand}(0, 1) < gmp$ then
        $\text{solution.p}[j] = \text{solution.p}[j] + (1 - \text{correctness}) \times \text{rand(solution.q}[j] - $\text{solution.p}[j], $1 - \text{solution.p}[j])$;
    end
end

Algorithm 6: Mutation operator for preference thresholds

input : the solutions to crossover: $parent\_1, parent\_2$; crossover probability; correctness of the solution; gene crossover probability ($gcp$)

if $\text{rand}(0, 1) < \text{crossover\_probability}$ then
    $\lambda = \text{rand}(0.1, 0.9)$;
    for $j = 1$ to $nb\_criteria$ do
        for $l = 1$ to $nb\_categories$ do
            if $\text{rand}(0, 1) < gcp$ then
                $\text{new\_sol\_1.profile}[j] = \lambda \times \text{parent\_1.profile}[j] + (1 - \lambda) \times \text{parent\_2.profile}[j]$;
                $\text{new\_sol\_2.profile}[j] = (1 - \lambda) \times \text{parent\_1.profile}[j] + \lambda \times \text{parent\_2.profile}[j]$;
            end
        end
    end
    if $\text{rand}(0, 1) < gcp$ then
        $\text{new\_sol\_1.w}[j] = \lambda \times \text{parent\_1.w}[j] + (1 - \lambda) \times \text{parent\_2.w}[j]$;
        $\text{new\_sol\_2.w}[j] = (1 - \lambda) \times \text{parent\_1.w}[j] + \lambda \times \text{parent\_2.w}[j]$;
    end
    if $\text{rand}(0, 1) < gcp$ then
        $\text{new\_sol\_1.q}[j] = \lambda \times \text{parent\_1.q}[j] + (1 - \lambda) \times \text{parent\_2.q}[j]$;
        $\text{new\_sol\_2.q}[j] = (1 - \lambda) \times \text{parent\_1.q}[j] + \lambda \times \text{parent\_2.q}[j]$;
    end
    if $\text{rand}(0, 1) < gcp$ then
        $\text{new\_sol\_1.p}[j] = \lambda \times \text{parent\_1.p}[j] + (1 - \lambda) \times \text{parent\_2.p}[j]$;
        $\text{new\_sol\_2.p}[j] = (1 - \lambda) \times \text{parent\_1.p}[j] + \lambda \times \text{parent\_2.p}[j]$;
    end
end

correct\_solution(\text{new\_sol\_1});
correct\_solution(\text{new\_sol\_2});

Algorithm 7: Crossover operator
chosen_solution = 0;
solution_1, solution_2 = randomly_chose_2_sol(population);
if rand(0, 1) < solution_1.objective/(solution_1.objective + solution_2.objective)
    then
        chosen_solution = solution_2;
    else
        chosen_solution = solution_1;
end

Algorithm 8: Selection operator


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