

# A Framework for Large-Scale Multiobjective Optimization Based on Problem Transformation

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**Abstract**—In this paper, we propose a new method for solving multiobjective optimization problems with a large number of decision variables. The proposed method called weighted optimization framework is intended to serve as a generic method that can be used with any population-based metaheuristic algorithm. After explaining some general issues of large-scale optimization, we introduce a problem transformation scheme that is used to reduce the dimensionality of the search space and search for improved solutions in the reduced subspace. This involves so-called weights that are applied to alter the decision variables and are also subject to optimization. Our method relies on grouping mechanisms and employs a population-based algorithm as an optimizer for both original variables and weight variables. Different grouping mechanisms and transformation functions within the framework are explained and their advantages and disadvantages are examined. Our experiments use test problems with 2–3 objectives 40–5000 variables. Using our approach on three well-known algorithms and comparing its performance with other large-scale optimizers, we show that our method can significantly outperform most existing methods in terms of solution quality as well as convergence rate on almost all tested problems for many-variable instances.

**Index Terms**—Large-scale optimization (LSO), many-variable optimization, metaheuristic framework, multiobjective optimization, variable grouping, weighting.

## I. INTRODUCTION

IN THE area of optimization, a growing interest in so-called large-scale optimization (LSO) can be observed [1]. This area of research deals with solving single-objective or multiobjective problems (MOPs) that include a large number of decision variables. The performance of classic metaheuristic algorithms often deteriorates when the dimensionality of

the decision space increases. When large numbers of variables are involved, algorithms are faced with very high-dimensional search spaces that are difficult to explore with limited resources.

Among large-scale problems, this paper focuses on multiobjective optimization problems that involve a large number of decision variables (many-variable problems). While single-objective LSO has already been extensively studied for many years [1], less work has been done to deal with multiobjective many-variable problems. Recently in 2015, a competition with single and multiple objectives based on a real-world EEG time series [2] data was held on the IEEE Congress on Evolutionary Computation (CEC) that involved up to 4864 decision variables. These kinds of applications emphasize the need for better algorithms to deal with large numbers of decision variables.

In this paper, we propose a new optimization framework called weighted optimization framework (WOF) which is based on a problem transformation, optimizing multiple weight vectors, and variable grouping. In our previous work [3], we briefly introduced the basic idea of WOF in a two-page short paper. This paper introduces and explains the WOF in detail by providing analysis based on theory and experimental data. The experiments performed in this paper use a larger set of benchmark functions and algorithms. Furthermore, we go into detail about different grouping mechanisms and transformation functions that can be used within the framework. The performance for different numbers of variables as well as further sensitivity analysis is provided. Finally, the results are compared with other LSO techniques.

The WOF is a new algorithm framework that enables population-based metaheuristics to solve many-variable problems more effectively and efficiently. In this paper, we show that the WOF can strongly enhance the performance of population-based metaheuristic algorithms to optimize many-variable multiobjective optimization problems. We provide an in-depth analysis of its functionality and give insight into its capabilities and limitations from a theoretical perspective. Additionally, we show that it can significantly outperform existing classical and large-scale algorithms on the tested multiobjective WFG [4], ZDT [5], DTLZ [6], and CEC2009 competition [7] problems with large numbers of variables.

This paper is organized as follows. In Section II, we give a short introduction about multiobjective optimization. Section III deals with occurring difficulties in LSO and gives a short overview of existing approaches for single-objective

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and multiobjective many-variable optimization. In Section IV, we propose the concept of the problem transformation, followed by explaining the details of the proposed optimization framework. Section V deals with different grouping mechanisms and transformation functions that are used in the framework. In our experiments in Section VI, we use the WOF framework on three well-known algorithms (NSGA-II [8], SMPSO [9], and GDE3 [10]) and compare their performances with each other and their original algorithm versions on a variety of well-known benchmark functions. Additionally, the multiobjective large-scale algorithms CCGDE3 [11], which is based on coevolution, and MOEA/DVA [12] are used for comparison. Finally, in Section VII, we conclude this paper.

## II. MULTIOBJECTIVE OPTIMIZATION

Many problems in the real world can be formulated as mathematical problems with certain goals (objectives) which have to be optimized. Additionally, some of these problems have constraints. If there are more than just one objective to optimize, such a problem is called an MOP. This can be formulated as follows:

$$\begin{aligned} Z: \quad & \min \quad \vec{f}(\vec{x}) = (f_1(\vec{x}), f_2(\vec{x}), \dots, f_m(\vec{x}))^T \\ & \text{s.t.} \quad \vec{x} \in \Omega \subseteq \mathbb{R}^n. \end{aligned} \quad (1)$$

This kind of MOP maps the decision space  $\Omega = \{\vec{x} \in \mathbb{R}^n \mid \vec{g}(\vec{x}) \leq 0\}$  of dimension  $n$  to the objective space  $M$  of dimension  $m$ . It is important to solve such a problem with or without constraints as accurate as possible. Since there are more than one objective, a single optimal solution can no longer be determined. Instead, modern problem solving methods concentrate on finding a Pareto-optimal solution set. However, because of the complexity of MOPs in real-world applications and limited computational resources, the true Pareto-front can hardly be analytically or exactly computed. Instead, heuristic algorithms are developed and used to find a nondominated solution set as good as possible to approximate the true Pareto-front.

## III. LARGE-SCALE OPTIMIZATION

LSO deals with optimizing problems that contain large numbers of variables (many-variable problems). Recently, a wide variety of single-objective many-variable optimizers have been proposed in the literature (see for instance the review by Mahdavi *et al.* [1]). Among others a competitive single-objective PSO algorithm [13] was developed in 2015 which showed good results for large-scale benchmark problems with up to 5000 decision variables. The majority of large-scale optimizers tend to use a concept called cooperative coevolution (CC). Since the pioneering work of Potter and De Jong [14], [15], the idea of CC has been used in various LSO algorithms. CC aims to optimize several independent populations, of which each holds a subset of the  $n$  decision variables. New solution candidates have to be formed by a combination of the variable values from different subcomponents, but genetic operators are used within one subcomponent only.

Several authors have used this concept for single-objective problems [16]–[21] and some work on multiple objectives also relied on coevolution [11], [22]. Regarding the distribution of the variables into subcomponents or groups, consideration for nonseparable problems was studied in [17], where the interaction of variables was taken into account by a learning mechanism for finding the optimal distribution of variables to the subcomponents. In [23], a mechanism called differential grouping (DG) was introduced to find improved distributions of the variables in single-objective CC algorithms. Other concepts and extensions to this approach for variable grouping have been proposed [24], [25]. In Section V, we go into detail on DG and other grouping mechanisms.

Our present work has been inspired by Yang *et al.* [16], where they applied a CC method to single-objective problems using two special features. One is a repeated grouping and regrouping of the variables into subcomponents in every iteration of the algorithm. The other is a weighting scheme to optimize a weight for each subcomponent, i.e., apply the same weight value (in terms of multiplication) to every variable in the same subcomponent. These weights were evolved for the best, worst and a random member of the population. The frequent (re)grouping in [16] was done randomly. The aspect that was a source of inspiration and that was adapted for the present work was their mechanism of optimizing a number of variables (in each subcomponent) at the same time by altering one weight value only. The same principle of using CC with weights has also been used in [18] and [19]. A mechanism for choosing appropriate lower and upper bounds of weights was proposed in [18]. More recently, Li and Yao [20] stated that using the weighting approach in CC is less effective than improving the frequent (re)grouping of variables. The above-mentioned CC-based studies used benchmark functions of up to 1000 decision variables.

In contrast to the above-mentioned and numerous other studies on large-scale single-objectives problems, less work has been conducted so far regarding multiobjective many-variable problems. In the area of multiobjective CC, good performance was reported in [11] for the ZDT [5] problems where CC was combined with a differential evolution algorithm called GDE3 [10]. They performed experiments with decision variables ranging from 200 up to 5000 and received good performance especially for the higher-dimensional problems. Nevertheless, the performance of the proposed method on more complicated benchmarks like the WFG toolkit or the DTLZ benchmarks has not been analyzed. Furthermore, although outperforming traditional methods by far, good approximations of the ZDT Pareto-front still needed a large number of function evaluations ranging between 150 000 to over 2 000 000 evaluations. An earlier approach by Iorio and Li [22] combined the concept of CC with the NSGA-II algorithm, but also focused on the ZDT problems for their analysis and tested rather small-dimensional instances of 10 and 30 variables.

For the real-world application in [2], good results compared to classical methods are shown in [26]–[28]. However, Zhang *et al.* [28] did only use the single-objective version of the problem, and Zhang *et al.* [26], [28] did not treat the

optimization problem in a black-box manner, as they used information obtained by analytical derivation of the objective functions. None of these three approaches used decomposition or CC approaches, and the performance on popular benchmarks as the ones frequently used by the community and in this paper have not been reported.

Two more very recent approaches to multiobjective LSO are introduced in [12] and [29]. In [12], a large-scale optimizer called MOEA/DVA is presented with the main focus on a new grouping mechanism, which aims to identify whether a decision variable contributes to convergence (i.e., distance to the Pareto-front), diversity, or both. Additionally, interactions between the variables are identified. Combining this information, a number of convergence-related, diversity-related as well as mixed groups are formed. The following optimization procedure of the MOEA/DVA focusses on the convergence/distance-variables first, before concentrating on the diversity. Ma *et al.* [12] reported a good performance for 200-variable instances of the UF1–UF6, UF10, ZDT4, and three-objective DTLZ1 and DTLZ3 benchmark functions. The grouping mechanism in this approach consumes a large amount of function evaluations prior to the start of the optimization, especially for large numbers of variables (refer to Section VI-E).

The most recent approach found to solve many-variable MOPs is called LMEA [29]. This method relies on a clustering approach to form groups of variables. It divides the variables into distance- and diversity-related groups. The experiments focused on many-variable instances and used selected five- and ten-objective DTLZ, WFG, and UF benchmark functions. LMEA showed good performance for 100, 500, and 1000 variable instances. It performed superior to the previously mentioned MOEA/DVA for the DTLZ1 and 2 (five-objective), DTLZ5–DTLZ7 and WFG3 problems, and worse on the UF9 and UF10 as well as the DTLZ4 (ten-objective) problems.

#### IV. PROPOSED METHOD

The aim of our proposed method is to reduce the dimensionality of optimization problems, so that existing algorithms are able to deal with them in a much more efficient manner. In contrast to the CC-based studies mentioned above, our proposed method is not using coevolution, but extending the concept of weighting the variables to multiple objectives instead. Similar to CC, our method distributes the decision variables to groups, but in contrast to CC it does not aim to optimize them independently. Our aim is to provide a multiobjective framework that makes use of the concept of optimizing smaller subproblems in a different way than current coevolutionary methods. Note that we do not intend to propose a new or improved grouping mechanism. The focus lies on proposing a framework that makes use of an arbitrary grouping mechanism and that is generic so that any grouping mechanism from the literature can be used. In this section, we first explain the concept of problem transformation and then continue with the proposed WOF algorithm that makes use of this transformation.

##### A. Problem Transformation

Let  $Z$  be a multiobjective optimization problem with  $n$  decision variables and  $m$  objectives as denoted earlier in (1). We

are interested in the set of Pareto-optimal solutions  $P := \{\vec{x}^*\}$  (Pareto-set) and the corresponding objective values  $\{f(\vec{x}^*)\}$  (Pareto-front). For any solution vector  $\vec{x}$ , we can write  $\vec{f}(\vec{x})$  as  $f(\psi(\vec{w}_l, \vec{x}))$ , with  $\vec{w}_l = (1, \dots, 1)$  a vector of ones and  $\psi$  the transformation function  $\psi(\vec{w}_l, \vec{x}) := (w_{l1}x_1, \dots, w_{ln}x_n)$ . To change the given solution we can now change the values of  $\vec{w}_l$ . In this way, instead of optimizing the vector  $\vec{x}$ , for any fixed real values of  $\vec{x}$  we can optimize a vector  $\vec{w}$  to approximate an optimal solution. It can easily be seen that by doing so, an optimal solution to the optimization problem can be found [see (3)]. For the optimal weights we obtain  $\vec{f}(\psi(\vec{w}^*, \vec{x})) = \vec{f}(\vec{x}^*)$  for a fixed  $\vec{x}$

$$\begin{aligned} \forall x_i : \exists w_i : w_i x_i &= x_i^* \\ &\Rightarrow \\ \forall \vec{x} \in \mathbb{R}^n : \exists \vec{w} : \psi(\vec{w}, \vec{x}) &= \vec{x}^*. \end{aligned} \quad (2)$$

By taking a fixed solution  $\vec{x}'$ , we can reformulate the original optimization problem  $Z$  to a new problem  $Z_{\vec{x}'}$  which optimizes the vector  $\vec{w}$  using  $\vec{x}'$  as an input parameter

$$\begin{aligned} Z_{\vec{x}'} : \min \quad & \vec{f}_{\vec{x}'}(\vec{w}) = (f_{1,\vec{x}'}(\vec{w}), \dots, f_{m,\vec{x}'}(\vec{w}))^T \\ \text{s.t.} \quad & \vec{w} \in \Phi \subseteq \mathbb{R}^n \\ & f_{o,\vec{x}'}(\vec{w}) = f_o(\psi(\vec{w}, \vec{x}')) \quad \forall o = 1, \dots, m \\ & \psi(\vec{w}, \vec{x}') := (w_1 x'_1, \dots, w_n x'_n). \end{aligned} \quad (3)$$

This new optimization problem has still the same number of decision variables as the original one. To reduce the dimensionality of the problem, we divide the original variables into  $\gamma$  groups ( $g_1, \dots, g_\gamma$ ), each of size  $l$  ( $\gamma \cdot l = n$ ). Furthermore, instead of assigning one  $w_i$  for each  $x_i$ , only one weight  $w_j$  is used for all variables within the same group  $g_j$  with  $l$  variables. Therefore, we reformulate the above-mentioned transformation function as

$$\psi(\vec{w}, \vec{x}) = (\underbrace{w_1 x_1, \dots, w_1 x_l}_{w_1}, \dots, \underbrace{w_\gamma x_{n-l+1}, \dots, w_\gamma x_n}_{w_\gamma}). \quad (4)$$

In this way, the vector  $\vec{w}$  has only a reduced size of  $\gamma < n$  (same as the number of groups), and therefore our new optimization problem  $Z_{\vec{x}'}$  has only  $\gamma$  decision variables. For an arbitrary but fixed solution  $\vec{x}'$ , we can now try to compute an approximation of the Pareto-set by optimizing this new optimization problem instead.

Up to this point there is a drawback to this approach, since by grouping the original variables together, their values cannot be changed independently of each other any more. This limits the reachable solutions in the original search space to a great extent. The searchable subspace of  $\Omega$ , which can be reached by optimizing  $Z_{\vec{x}'}$ , is defined mainly by three choices.

- 1) The choice of  $\vec{x}'$ .
- 2) The choice of the grouping scheme, i.e., the decision of which variables are put together in the same group.
- 3) The choice of the transformation function  $\psi(\cdot)$ .

In the remainder of this paper, we will explain some possible approaches to each of these three choices in further detail. First, an appropriate choice of  $\vec{x}'$  is crucial for the success of the optimization since it defines a “direction” of the future



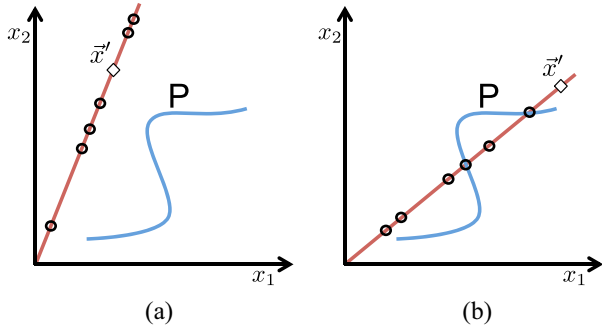


Fig. 1. Examples of variable grouping and a choice of  $\vec{x}'$  in the decision space. Solutions on the red straight line can be obtained after  $x_1$  and  $x_2$  are grouped together.  $P$  denotes the Pareto-optimal set in the decision space. Note that even the suboptimal choice on the left might be beneficial to the search, as it allows to advance closer to certain parts of the Pareto-set. (a) “Suboptimal” choice of  $\vec{x}'$ . (b) “Optimal” choice of  $\vec{x}'$ .

search. This shall be visualized by a small example. Consider a simple optimization problem with just two decision variables  $x_1$  and  $x_2$ . In Fig. 1, we show the search space of this problem, as well as a hypothetical (Pareto-)optimal set denoted as  $P$ . When optimizing the original problem  $Z$ ,  $x_1$  and  $x_2$  are changed independently of each other and all of the solutions in the search space can be found. Therefore, a good approximation of  $P$  can be obtained. The effect of grouping  $x_1$  and  $x_2$  together and forming a new problem  $Z_{\vec{x}'}$  will result in the optimization of just one single variable  $w_1$ . This limits the reachable solutions to a 1-D subspace (denoted as a red line from  $\vec{x}'$  to the origin in Fig. 1). Comparing the scenes of the Fig. 1(a) and (b), we see that some optimal solutions can or cannot be reached depending on the choice of  $\vec{x}'$ . Carrying this knowledge to higher-dimensional problems, every group of variables forms a subset of configurations that are reachable by the new optimization process. However, note that even the “suboptimal” choice of  $\vec{x}'$  in Fig. 1(a) might be beneficial to the overall search, as it allows the optimization process to advance closer to certain parts of the Pareto-set.

In a similar way, this accessible subspace is influenced by the grouping scheme and transformation function. The grouping of the variables has, similar to the location of  $\vec{x}'$ , an influence on the subspace of  $\mathbb{R}^n$  in which the search will take place. The transformation function on the other hand mainly describes how the movement takes place inside this space, i.e., in which way the locations of the newly created solutions change. Additionally, it can affect the shape of the lower-dimensional space (i.e., a line like in Fig. 1). The transformation of the problem, i.e., the definition of the searchable subspace that is accessible from a fixed solution, could also be seen as a kind of “neighborhood” of the original solution  $\vec{x}'$ , in the sense that only certain directions of change in a solution are possible starting from  $\vec{x}'$ . Consequently, different transformation functions [other than in (4)] produce different neighborhoods. However, in contrast to neighborhoods in local search methods, the solutions created in these neighborhoods in WOF might not necessarily have a locality around the original solution. In fact, the opposite might be the case, as the steps made in the decision space might be very large depending on the weight values.

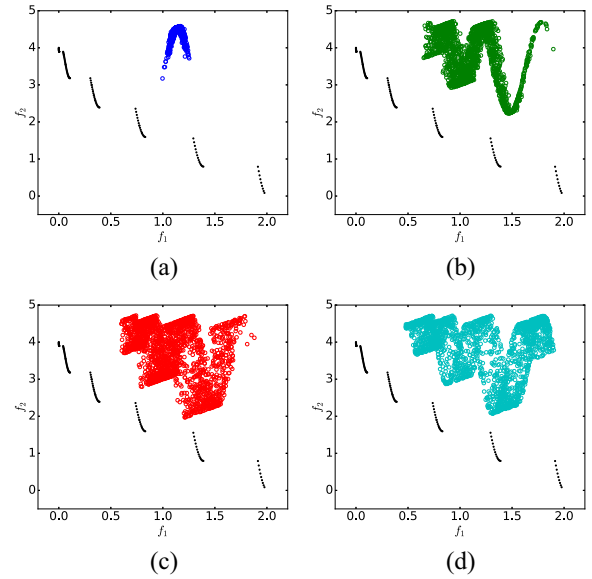


Fig. 2. Pareto-front (small black dots) and created solutions for the WFG2 problem by classic EA methods and different WOF groupings. (a) Crossover and mutation. (b) Random grouping. (c) Ordered grouping. (d) DG.

Although the explained problem transformation has disadvantages in preserving diversity and therefore in the ability to approximate the whole Pareto-optimal set, it must be mentioned that it can, on the other hand, have positive effects on the convergence speed, if a good choice of  $\vec{x}'$  is made and appropriate grouping schemes and transformation functions are used. Different grouping schemes and transformation function will be explained in Section V. Here we want to point out that once good choices for these parameters are found, the new smaller search space can be searched much faster and more thoroughly.

To examine the grouping and transformation process and its implications in further detail, in Figs. 2–4 we show the effect of this approach exemplary on the three test problems WFG2, UF2 (from the CEC2009 competition) and ZDT1 with  $n = 1000$  decision variables. All plots show 2000 solutions created from a fixed randomly created solution by different methods. Figs. 2(a)–4(a) show the solutions obtained by a standard SBX-crossover and polynomial mutation. For each of the 2000 solutions shown in Figs. 2(a)–4(a), another solution was randomly created and used for crossover. The mutation and crossover parameters were the same as reported in Section VI-A. Using the same solution as in Figs. 2(a)–4(a), instead of applying crossover and mutation, the proposed problem transformation was carried out with different grouping mechanisms in Figs. 2(b)–(d)–4(b)–(d) (for details on these mechanisms see Section V-A). For each of these transformed problems, 2000 random solutions, i.e., weight vectors were created.

We can see in these figures that for all three problems the solutions obtained by the problem transformation differ to a great extent from the ones accessible by traditional evolutionary operators. In case of the WFG2 problem, the reachable solutions offer by far more diversity as well as closeness to the Pareto-front. If the transformed problem is optimized to obtain

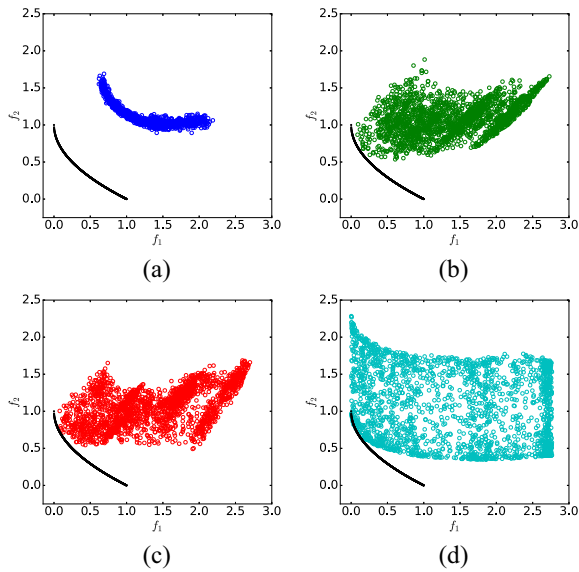


Fig. 3. Pareto-front (small black dots) and created solutions for the CEC2009 UF2 problem by classic EA methods and different WOF groupings. (a) Crossover and mutation. (b) Random grouping. (c) Ordered grouping. (d) DG.

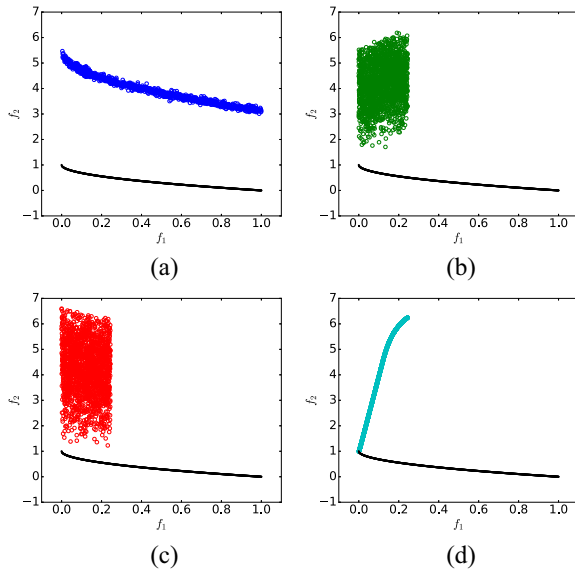


Fig. 4. Pareto-front (small black dots) and created solutions for the ZDT1 problem by classic EA methods and different WOF groupings. (a) Crossover and mutation. (b) Random grouping. (c) Ordered grouping. (d) DG.

the best of the solutions seen in Fig. 2(b)–(d), an immense speed up of the search can be expected.

Almost the same situation occurs when transforming the UF2 problem in Fig. 3. The effects of the grouping mechanisms differ slightly; however, all these transformed problems offer the possibility to reach a set as diverse as the one reached by classical operators. Moreover, the solutions obtainable lie very close to the Pareto-front.

The situation in Fig. 4 when transforming the ZDT1 problem differs significantly from the ones in Figs. 2 and 3. In Fig. 4, we see the effect of losing diversity that was described

previously. While the classical crossover and mutation operators do offer a great extend of diversity, mostly because these solutions are based on crossover with other solutions, the transformed random solutions are limited to a part of the objective space as seen in Fig. 4(b)–(d). The area to which the search is limited is defined by the chosen  $\vec{x}'$  in this case, which happened to be on the “left side” of the Pareto-front. Therefore, optimizing only the transformed problem might lead to fast convergence in Fig. 4(d), but might suffer from a very low diversity.

### B. Weighted Optimization Framework

In this section we describe our proposed approach in detail. The WOF is designed as a generic meta-heuristic, which can be incorporated in an arbitrary population-based optimization mechanism. The choice of this mechanism depends on the preferences of the user or the properties of the problem. This optimization algorithm will be used for optimizing two different problems  $Z$  and  $Z_{\vec{x}'}$  in turns. It is also possible to use two different optimization algorithms for the two problems. This might be advisable as the two problems differ largely in size. For instance, the smaller transformed problem could be thoroughly searched with a conventional method, while the original, large-scale problem can be searched with a CC-based method. This paper, however, concentrates on employing only one optimizer for both problems. The use of two different methods for the two optimization phases is left for future work.

As we have seen in the previous subsection, the transformed problem  $Z_{\vec{x}'}$  has the drawback of limiting the search space and the advantage of searching a smaller space more thoroughly, which might result in a faster convergence but less diversity. On the other hand, the original problem  $Z$  can reach all possible solutions but might incorporate very slow convergence in a large-dimensional space. To utilize the synergy of these two formulations, WOF alternates two different phases of optimization: a *normal* optimization step and a *weighting* optimization step. The outline of our proposed method is shown in Fig. 5 and more detailed steps are given as pseudocode in Algorithms 1–3.

The WOF algorithm receives an optimization problem  $Z$ , a population-based optimization algorithm  $A$ , a grouping mechanism  $G$  (see Section V-A) as well as a transformation function  $\psi$  (see Section V-B) as its input. The starting point is an initial random population for the problem  $Z$ . In every iteration of the main loop (i.e., lines 3–10 in Algorithm 1), first the normal optimization step is carried out, where the original problem is optimized for a fixed number  $t_1$  of function evaluations (line 4 in Algorithm 1). The only special requirement here is that the algorithm needs to start with a given population instead of creating a new random one.

Next, in the weighting optimization step (lines 6–8 in Algorithm 1) the transformation and optimization of weights are done  $q$  times to preserve diversity in the population (details below in Section IV-C). For this,  $q$  different solutions  $\vec{x}'_k$  ( $k = 1, 2, \dots, q$ ) are drawn from the current population (line 5). For every one of them, the weighting optimization is carried out as shown in Algorithm 2. The problem is first

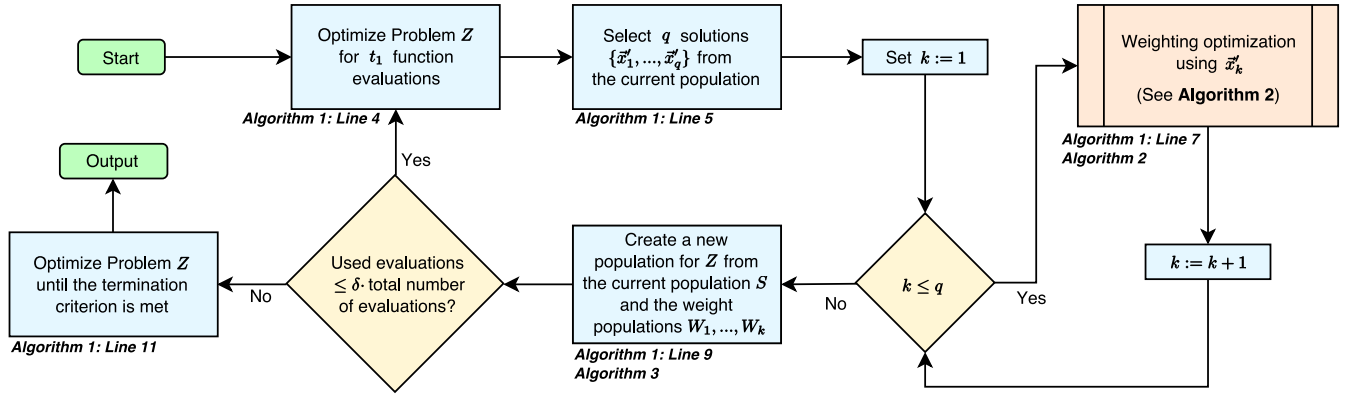


Fig. 5. Outline of the WOF.

**Algorithm 1** WOF( $Z, A, G, \psi$ )

**Input:** Problem  $Z$ , Optimization Algorithm  $A$ , Grouping Mechanism  $G$ , Transformation Function  $\psi$

**Output:** Solution population  $S$

- 1: Initialization
- 2:  $S \leftarrow$  Random initial population for  $Z$
- 3: **repeat**
- 4:  $S \leftarrow A(Z, S, t_1)$  //Optimize  $Z$  with Algorithm  $A$  for  $t_1$  evaluations, using  $S$  as a starting population.
- 5:  $\{\vec{x}'_1, \dots, \vec{x}'_q\} \leftarrow$  Selection of  $q$  solutions based on Crowding Distance from the first nondominated front of  $S$
- 6: **for**  $k = 1$  **to**  $q$  **do**
- 7:  $W_k \leftarrow$  WeightingOptimization( $\vec{x}'_k, Z, A, G, \psi$ ) //Algorithm 2
- 8: **end for**
- 9:  $S \leftarrow$  updatePopulation( $W_1, \dots, W_q, S$ ) //Algorithm 3
- 10: **until**  $\delta \cdot \text{total\#Evaluations}$  used
- 11:  $S \leftarrow A(Z, S)$  //Optimize  $Z$  with Algorithm  $A$ , using  $S$  as starting population, until all evaluations are used.
- 12: **return** FirstNonDominatedFront( $S$ )

**Algorithm 2** WeightingOptimization( $\vec{x}'_k, Z, A, G, \psi$ )

**Input:** Solution  $\vec{x}'_k$ , Problem  $Z$ , Optimization Algorithm  $A$ , Grouping Mechanism  $G$ , Transformation Function  $\psi$

**Output:** Population of weights  $W_k$

- 1: Initialization
- 2: Divide  $n$  variables into  $\gamma$  groups //See **Subsection V-A**
- 3:  $Z_{\vec{x}'}^k \leftarrow$  Build a transformed problem with  $\gamma$  decision variables (weights) from  $Z, \vec{x}'_k, G$  and  $\psi$
- 4:  $W_k \leftarrow$  Random population of weights for  $Z_{\vec{x}'}^k$
- 5:  $W_k \leftarrow A(Z_{\vec{x}'}^k, W_k, t_2)$  //Optimize  $Z_{\vec{x}'}^k$  with Algorithm  $A$  for  $t_2$  evaluations, using  $W_k$  as a starting population.
- 6: **return**  $W_k$

**Algorithm 3** UpdatePopulation( $W_1, \dots, W_q, S$ )

**Input:** Weight populations  $W_1, \dots, W_q$ , Solution population  $S$

**Output:** Solution population  $S$

- 1: **for**  $k = 1$  **to**  $q$  **do**
- 2:  $w_k \leftarrow$  Select one individual from  $W_k$
- 3:  $S'_k \leftarrow$  Apply  $w_k$  to population  $S$
- 4: **end for**
- 5:  $S \leftarrow$  Perform non-dominated sorting on  $S \cup \{S'_k\}_{k=1, \dots, q}$
- 6: **return**  $S$

transformed as described above, resulting in different transformed problems  $Z_{\vec{x}'}^k$ . At this point, a transformation function has to be applied, and as mentioned earlier, different functions can be used. It is also possible to choose different transformation functions for each of the  $q$  transformed problems. The optimization algorithm is then used to optimize a population  $W_k$  of randomly created solutions (weights) for each of these  $Z_{\vec{x}'}^k$ . The result of this low-dimensional optimization step is a population  $W_k$  of weights that optimizes the objective function values based on the values of the originally chosen solution  $\vec{x}'_k$ . This whole process of problem transformation and optimization is carried out  $q$  times, giving us a set of  $q$  weight populations  $\{W_1, \dots, W_q\}$ .

These obtained weights are then assigned to the original solution population (line 9 in Algorithm 1) in the following way, as also shown in detail in Algorithm 3. In order to save computational resources, from every  $W_k$ , we select one member (with the largest crowding distance from the first non-dominated front) and apply its values to every solution in  $S$ . By

doing so, we obtain  $q$  sets of new solutions  $S'_k$  ( $k = 1, \dots, q$ ). To update the original population, we combine the obtained new solution sets with  $S$  and perform a nondominated sorting mechanism as used in NSGA-II on the set  $S \cup \{S'_k\}_{k=1, \dots, q}$ . Duplicate solutions found in this union set are removed prior to the nondominated sorting, as they would deteriorate diversity. The nondominated sorting procedure in this step also ensures that worse solutions found in the current iteration (for example due to a suboptimal choice of one of the  $\vec{x}'_k$ ) will not be further regarded in the upcoming steps. After that, the main loop starts again and we return to a normal optimization step to alter the variable values independently of each other.

In addition to applying one individual of each  $W_k$  to the population  $S$ , it is also possible to add all members of each  $W_k$  to the population update process. These solutions have already been evaluated in the optimization of  $Z_{\vec{x}'}^k$  and can therefore be included in the process. Depending on the population size used

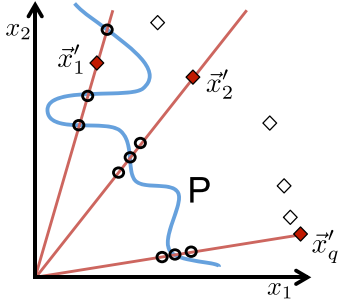


Fig. 6.  $q$  optimization steps in WOF. Diamonds: original population. Circles: achieved solutions of the  $q$  transformed problems.

when optimizing  $Z_{\vec{x}}^k$ , this might improve the results slightly in favor of a slightly longer computation time. However, previous experiments showed, that by doing so the improvement in performance is, in most cases, negligible.

The alternation of the main loop of WOF (lines 3–10 in Algorithm 1) is only repeated until the first termination condition is met. More precisely, the optimization of variables and weights will only take place during the first  $\delta \cdot total\#Evaluations$ , where  $\delta \in [0, 1]$  defines how many of the total function evaluations are spent for this phase. After that, the algorithm will use the normal optimization without weights for the second half of the optimization process (line 11 in Algorithm 1).

The reason for this alternation is that the transformation of the problem might result in a directed search where only a certain area of the Pareto-front can be reached. Early experiments showed that using the weight optimization throughout the whole search process results in an overall loss of diversity. In general, the efficiency of WOF is most probably higher in the initial phase of an optimization process than in the final phase. Toward the end of the search, the Pareto-front might already be approximated to a certain extent. When WOF is used in this situation, the grouped alternation might deteriorate optimal values of one variable in favor of another. In this phase of the search, classical (ungrouped) evolutionary operators can make finer adjustments to the decision variable values independently of each other and therefore slowly approach the optimal values for each single variable.

### C. Parameter $q$ and Choice of $\vec{x}'$

Now we take a look at how to choose the  $\vec{x}'$  for the weighted optimization. In the case of just one objective function the selection is rather straightforward, as we can determine the *best* and *worst* solutions. Unfortunately we do not have a total order on the solutions in multiobjective optimization, so choosing one  $\vec{x}'$  might result in a great acceleration of the optimization process toward (a part of) the Pareto-front, but in exchange, we might lose the diversity of the population [see Fig. 4(d)]. To deal with this problem, we perform multiple independent weighting optimizations using different  $\vec{x}'_k$ ,  $k = 1, \dots, q$ . It might be advisable to specify  $q$  as  $m$  or larger (where  $m$  is the number of objectives), and decide to make the selection of  $\vec{x}'$  based on a diversity indicator. In this way we can choose multiple solutions that lie “far” from each other in the objective space, to hopefully cover different

areas of the decision space with the weighting optimization as shown in Fig. 6. In this paper, we select the best  $q$  solutions by applying the crowding distance metric to the first nondominated front of the current population of the original problem. It is also possible to use other selection criteria in this step. In our case, we have obtained better results using crowding distance compared to a random selection (refer to Tables 17–19 and Figs. 42–45 in the supplementary material). For problems with many objectives it may be promising to use reference-line-based approaches.

## V. GROUPING AND TRANSFORMATION FUNCTIONS

In this section, we will describe four different mechanisms for grouping variables and three transformation functions that will be used later in the experiments.

### A. Grouping Mechanisms

Grouping strategies usually aim to put those variables into the same group that interact strongly with each other (in nonseparable optimization problems). Here we briefly explain different grouping methods that are used in this paper to examine the influence of a good grouping strategy. While the first three methods are rather simple and do not use any information about the objective functions, the DG incorporates an intelligent mechanism based on problem analysis. We use this method for comparison, although it has been developed for single-objective optimization.

1) *Random Grouping*: Random grouping forms a fixed number  $\gamma$  of equal-sized groups and assigns each variable randomly to each of these groups.

2) *Linear Grouping*: Linear grouping assigns all  $n$  variables to a fixed number  $\gamma$  of groups in natural order. This means, the first  $(n/\gamma)$  variables of the optimization problem are assigned to the first group and so on. This corresponds to the situation seen in (4).

3) *Ordered Grouping*: The ordered grouping mechanism ranks the decision variables of the selected solution by their absolute values. This means that all variables are sorted based on their current absolute values and the  $(n/\gamma)$  variables with the smallest values are assigned to the first group, the next  $(n/\gamma)$  to the second group, and so forth.

4) *Differential Grouping*: DG was developed in 2014 by Omidvar *et al.* [23] and used in single-objective optimization with CC. It aims to detect variable interaction prior to the optimization of the problem. The number of groups as well as their sizes are set automatically by the DG algorithm. In short, DG compares the amount of change in the objective function in reaction to a change in a variable  $x_i$  before and after another variable  $x_h$  is changed. DG answers the following question: *when changing the value of  $x_i$ , does the amount of change in  $f(\vec{x})$  remains the same regardless of the value of another variable  $x_h$ ?* If this is true, the variables  $x_i$  and  $x_h$  seem not to interact with each other, so they can be separated into different groups. Else, they seem to interact, so they are assigned to the same group. Two drawbacks must be mentioned when using this approach here.

1) The DG algorithm can become computationally expensive as analyzed in [23]. Assuming the problem contains



a number  $\gamma = (n/l)$  evenly sized groups with  $l$  variables each, the number of function evaluations consumed is in  $O(n^2/l)$ . This means for a fully separable problem using  $n = 2000$  decision variables, DG requires  $n^2 = 4\,000\,000$  function evaluations to perform the grouping. Given that our experiments in Section VI only use 100 000 function evaluations for the whole optimization process, this is little practicable.

- 2) As mentioned, the DG algorithm was developed for single-objective problems. It was not designed to take multiple objective functions into account and, as a result, is not directly applicable to MOPs. To make it applicable to multiobjective optimization, in this paper only one of the objective functions (the first one) is considered in the DG algorithm. The implications and possibly different outcomes when using other than the first objective function might be studied in a future work.

The four grouping mechanisms differ in the required amount of computational effort in each generation. The linear grouping does not change during the process and therefore does not require recomputation in each generation. The same holds for DG, which is precomputed before the optimization starts and not changed during the search. On the other hand, the random grouping and the ordered grouping mechanisms are updated every time a problem transformation is carried out. In the former, a new random assignment to the  $\gamma$  groups is computed, in the latter the variables of each chosen  $x'_k$  are ordered in each problem transformation step.

### B. Transformation Functions

In the following, we will describe three different transformation functions  $\psi(\vec{w}, \vec{x}')$  and examine their advantages and disadvantages. We assume the variables were divided into  $\gamma$  groups  $g_1, \dots, g_\gamma$  prior to the transformation. Let  $g(x_i) = g_j$  be the respective group  $x_i$  is assigned to (i.e., the set of variables in the same group as  $x_i$ ) and  $w_j$  the respective weight used to transform the values of  $g_j$ .

1)  $\psi_1$ —*Product Transformation*: In Section IV-A, we used a basic function  $\psi(\vec{w}, \vec{x}') := (w_1 x'_1, \dots, w_\gamma x'_\gamma)$ . Broken down to a single element, this equation results in the simple expression  $x_{i,\text{new}} = w_j \cdot x_{i,\text{old}}$  for each variable. Therefore, the first transformation function is as follows:

$$\begin{aligned} x_{i,\text{new}} &= w_j \cdot x_{i,\text{old}} \\ w_j &\in [0, 2]. \end{aligned} \quad (5)$$

A disadvantage of  $\psi_1$  is that we do not expect it to deal well with variable domains that involve both positive and negative values (since a multiplication with positive weights can only obtain positive or negative new values, respectively). Additionally, the progress that can be made by altering one weight variable  $w_j$  is determined by the absolute value of  $x_i$ . The genetic operators used to alter the  $w_j$  are naturally limited in the amount of change per iteration, while at the same time we also need to apply some sort of domain to the  $w_j$  during the optimization process. That means, if the variable  $x_i$  has a domain of  $x_i \in [0, 10]$ , and the variable  $w_j$  has a domain of  $w_j \in [0, 2]$ , an absolute value of  $x_i = 0.1$  will

give the optimization algorithm only a maximum chance to explore the original search space in the interval  $[0, 0.2]$ , while a value of  $x_i = 8$  will result in a search that covers at least the whole domain of  $x_i$ . Since we do not know the optimal variable values beforehand, the search should not be dependent on absolute values. Therefore, we propose the following transformation function.

2)  $\psi_2$ —*p-Value Transformation*: This method is intended to uncouple the relationship between absolute variable values and the reachable domain space of the original variable

$$\begin{aligned} x_{i,\text{new}} &= x_{i,\text{old}} + p \cdot (x_{i,\text{max}} - x_{i,\text{min}}) \cdot (w_j - 1.0) \\ w_j &\in [0, 2] \\ p &\in [0, 1]. \end{aligned} \quad (6)$$

Using this function, the value of  $x_i$  is always altered within a certain range (defined by  $p$ ) around its original value, where  $x_{i,\text{min}}$  and  $x_{i,\text{max}}$  are the lower and upper bounds of the variable  $x_i$ . The values of the  $w_j$  are bound to the interval  $[0, 2]$  to ensure that variable values during optimization are always positive. As can be seen in the first line of the equation, these values are translated into  $[-1, 1]$ , which describes a change of values in one or the other direction around the original value. The actual maximum and minimum amounts of change are set by the parameter  $p$ , so that the domain of the  $w_j$  does not have to be altered. For instance, setting  $p = 0.2$ , the value of  $x_i$  is altered by 20% of the width of the variable's domain, centered around the original value  $x_{i,\text{old}}$ .

3)  $\psi_3$ —*Interval-Intersection Transformation*: This function follows an approach suggested in [18] for single-objective optimization. We include it in this paper especially for the comparison on the ZDT functions. This is because the previous two functions need a kind of *repair-mechanism* in case the result of the transformation function exceeds a variables domain. In this case, both  $\psi_1$  and  $\psi_2$  set the value of the variable to the respective boundary. However, this might be of advantage for problems like the ZDT benchmarks, since their optimal values are extremal values and can be obtained by such repairing. To provide a fair comparison with other algorithms on the ZDT problems in Section VI-D, we use the following transformation function  $\psi_3$ :

$$\begin{aligned} x_{i,\text{new}} &= w_j \cdot x_{i,\text{old}} \\ w_j &\in \left[ \min_{x_h \in g(x_i)} (x_{h,\text{min}}/x_h), \max_{x_h \in g(x_i)} (x_{h,\text{max}}/x_h) \right] \end{aligned} \quad (7)$$

where  $x_{h,\text{min}}$  and  $x_{h,\text{max}}$  are the lower and upper bounds of the variable  $x_h$ . The application of the weight value follows the same multiplication equation as in  $\psi_1$ , only the domains of the weights for each group are changed. For each variable in a group, the maximum (minimum) weight value that can be used without exceeding the domain borders is calculated. The upper- and lower-bounds of the variables  $w_j$  are set based on the highest (lowest) of these possible values over all variables in the respective group. This interval  $[w_{j,\text{min}}, w_{j,\text{max}}]$  can be seen as the intersection interval of all possible minimum and maximum values defined by each variable in the group.



## VI. EVALUATION

The benchmark problems used in this paper are WFG1–WFG9 [4], DTLZ1–DTLZ7 [6], ZDT1–ZDT4 and ZDT6 [5], and the ten unconstrained problems of the CEC2009 competition on multiobjective optimization (denoted as UF1–UF10) [7]. With the exception of the CCGDE3 algorithm, we rely on the jMetal framework for Java, version 4.5 [30] for the implementation of the algorithms. The source codes of the mentioned problems are included in the framework. For the comparison with the CCGDE3 algorithm, we obtained the original code from the website of the authors.<sup>1</sup> The C++ implementations of the UF problems were taken from the original competitions resources. The original C++ implementation of the WFG group was used, and the DTLZ C++ implementation were taken from the C++ adaptation of jMetal.

The algorithms used in this evaluation are as follows.

- 1) SMPSO, which represents a state-of-the-art swarm optimizer.
- 2) NSGA-II, which is a widely known multiobjective evolutionary algorithm.
- 3) GDE3, which uses differential evolution.
- 4) WOF-SMPSO, WOF-NSGA-II, and WOF-GDE3, which are WOF-enhanced versions of the previous ones, i.e., the respective algorithm (SMPSO, NSGA-II, and GDE3) is used as an optimizer within our proposed framework to optimize the weights and variables.
- 5) CCGDE3 is used as another large-scale optimizer. Published in 2013, it is a recent algorithm designed specifically for multiobjective many-variable problems.

The performance indicator used in this paper is the hypervolume indicator [31]. It can be used to measure both convergence and diversity of an obtained solution set. When reporting hypervolume values in this paper, relative values are used, i.e., the obtained hypervolume divided by the hypervolume of the true Pareto-front. The reference point for calculating the hypervolume in all instances is obtained from a sample of the true Pareto-front of each problem by multiplying the nadir-point of the sample by 2.0 in each dimension. For each algorithm and each tested problem, we perform 51 independent runs. All experiments are conducted with two and three objectives wherever possible, and use decision variables from 40 up to 5000 depending on the respective experiment. For testing the statistical significance of the differences between indicator values (which are not likely to follow a normal distribution) we use the Mann-Whitney U statistical test. We indicate a difference as significant for a value of  $p < 0.01$ .

### A. General Parameter Settings

Many of the experiments in the following sections will use identical parameter settings. In the following sections, only settings that differ from those given here will be explained.

The maximum number of function evaluations is set to 100 000 in all experiments. In all WOF algorithms, the number  $t_1$  of evaluations for each optimization of the original problem

$Z$  is set to 1000, and the number of evaluations for the transformed problem  $t_2$  is 500. The parameter  $g$ , the number of repetitions of the weight optimization (therefore also the number of chosen solutions  $\vec{x}'$ ) is set to 3, and the number of groups ( $\gamma$ ) is set to 4 in the WOF algorithms and the CCGDE3. The ordered grouping mechanism is used in all experiments and the transformation function is  $\psi_2$  with  $p = 0.2$ . The parameter  $\delta$ , which defines the evaluations spent for the first phase (lines 3–10 in Algorithm 1) is set to 0.5. A dynamic version where the termination of the first phase is linked to the progress of the search might be possible too, however, this task is left for future work. The sensitivity of WOF to different values of  $\delta$  is reported later in Section VI-G.

SMPSO and NSGA-II use polynomial mutation with a probability of  $1/n$  (with  $n$  being the number of decision variables) and a distribution index of 20.0. The NSGA-II and its WOF-enhanced version use an SBX crossover with a crossover probability of 0.9 and a distribution index of 20.0. In GDE3, CCGDE3, and WOF-GDE3, the parameters  $F$  and  $CR$  are both set to 0.5.

The size of the populations as well as the archives is set to 100 for all algorithms. In WOF, the optimization of the weights takes place with a population size of 10 due to the reduced size of the search space. The CCGDE3 uses an internal population size of 40, but creates a larger population with up to 160 solutions when the algorithm returns. We wanted to keep the parameters of the CCGDE3 the same as in the original paper, so the same setting is used here, with the exception that only the best 100 solutions are returned to ensure fairness to the other methods. The number of generations per single group in CCGDE3 also follows the original publication and is set to 1.

### B. Performance on Various Benchmarks

In this part, we will compare the performance of all algorithms on the mentioned benchmarks. All test problem instances use  $n = 1000$  decision variables in this experiment. For the WFG problems, they are split into 250 position-related and 750 distance-related variables.

The median relative hypervolume values and interquartile ranges (IQRs) can be seen in Table I. Best performance is indicated by a bold font. An asterisk is used to indicate whether the result is significantly worse than the respective best algorithm's performance for each problem. For readability, values equal to 0.0 are displayed as dashes.

We see that the WOF algorithms perform on average very well. For almost all of the tested problems, at least one version of the WOF algorithms performs significantly better than all the non-WOF methods. In the following, some interesting observations will be pointed out for the different problem families.

1) *WFG*: In the WFG problems none of the classic algorithms nor CCGDE3 can obtain a best hypervolume value. Moreover, all of the results are significantly worse than the respective best algorithm. In Table I, the best results are obtained by WOF-SMPSO for almost all test problems except for WFG1 with two and three objectives and WFG5 with two objectives (for which no significant difference between

<sup>1</sup><https://www.cs.cinvestav.mx/ EVOCINV/software/CCLSMO/CCLSMO.html>

TABLE I  
 MEDIAN RELATIVE HYPERVOLUME VALUES OF VARIOUS BENCHMARK PROBLEMS FOR  $n = 1000$  DECISION VARIABLES. BEST PERFORMANCE IS SHOWN IN BOLD FONT. AN ASTERISK INDICATES STATISTICAL SIGNIFICANCE COMPARED TO THE RESPECTIVE BEST METHOD

m		CCGDE3	GDE3	NSGAI	SMPSO	WOF-GDE3	WOF-NSGAI	WOF-SMPSO
2	DTLZ1	— (—) *	— (—) *	— (—) *	0.642949 (—)	— (—) *	— (—) *	0.642949 (—)
2	DTLZ2	— (—) *	— (—) *	— (—) *	<b>0.998909</b> (5.10E-5)	0.995145 (1.49E-3) *	0.997587 (4.00E-4) *	0.998694 (1.26E-4) *
2	DTLZ3	— (—) *	— (—) *	— (—) *	0.520072 (—)	— (—) *	— (—) *	0.520072 (—)
2	DTLZ4	— (—) *	— (—) *	— (—) *	<b>0.998900</b> (5.00E-5)	0.987029 (1.94E-2) *	0.996243 (7.80E-4) *	0.998700 (1.49E-4) *
2	DTLZ5	— (—) *	— (—) *	— (—) *	<b>0.999348</b> (5.83E-5)	0.995736 (2.90E-3) *	0.998003 (4.86E-4) *	0.999100 (1.36E-4) *
2	DTLZ6	— (—) *	— (—) *	— (—) *	— (—) *	0.999011 (1.45E-4) *	<b>0.999150</b> (6.17E-5)	0.999149 (3.75E-1)
2	DTLZ7	<b>0.698002</b> (7.78E-3) *	<b>0.380034</b> (1.40E-2) *	<b>0.442085</b> (2.36E-2) *	<b>0.070975</b> (5.03E-2) *	<b>0.798233</b> (1.01E-5) *	<b>0.798236</b> (1.51E-5) *	<b>0.798270</b> (6.21E-6)
2	WFG1	0.368023 (9.71E-3) *	0.575510 (8.54E-3) *	0.314626 (2.34E-3) *	0.590658 (7.06E-3) *	<b>0.652464</b> (1.66E-2) *	0.646211 (4.40E-3) *	0.645470 (1.11E-2) *
2	WFG2	0.619606 (5.85E-3) *	0.688903 (4.22E-3) *	0.667313 (9.52E-3) *	0.736841 (2.53E-2) *	0.862312 (3.16E-2) *	0.800874 (1.41E-2) *	<b>0.983370</b> (6.56E-3) *
2	WFG3	0.543414 (6.03E-3) *	0.683368 (5.56E-3) *	0.612655 (7.43E-3) *	0.594265 (2.81E-2) *	0.807946 (8.23E-3) *	0.807269 (6.12E-3) *	<b>0.855820</b> (4.67E-3) *
2	WFG4	0.507776 (6.72E-3) *	0.630822 (1.46E-2) *	0.621814 (9.71E-3) *	0.843217 (4.46E-3) *	0.870188 (2.07E-2) *	0.856543 (1.47E-2) *	<b>0.975442</b> (1.26E-2) *
2	WFG5	0.541706 (6.72E-3) *	0.668053 (2.06E-2) *	0.587745 (1.11E-2) *	0.803472 (4.10E-3) *	0.940453 (7.84E-2) *	0.878879 (9.23E-2) *	<b>0.946467</b> (3.20E-2) *
2	WFG6	0.400922 (8.92E-3) *	0.495129 (1.14E-2) *	0.576546 (8.92E-3) *	0.969041 (4.28E-3) *	0.906795 (4.52E-2) *	0.834530 (5.97E-2) *	<b>0.998855</b> (2.55E-4) *
2	WFG7	0.524232 (1.18E-2) *	0.696964 (1.52E-2) *	0.648534 (8.83E-3) *	0.697714 (1.82E-2) *	0.857582 (4.17E-3) *	0.862129 (5.15E-3) *	<b>0.962463</b> (1.35E-2) *
2	WFG8	0.466575 (5.19E-3) *	0.667567 (6.39E-3) *	0.551133 (1.01E-2) *	0.515858 (7.58E-4) *	0.810636 (1.97E-2) *	0.815991 (2.46E-2) *	<b>0.885010</b> (1.91E-2) *
2	WFG9	0.473511 (1.08E-2) *	0.511297 (9.30E-3) *	0.621036 (3.40E-2) *	0.894743 (9.31E-3) *	0.935509 (1.96E-2) *	0.931595 (2.27E-2) *	<b>0.962831</b> (1.47E-2) *
2	UF1	0.768177 (2.06E-2) *	0.673944 (3.26E-2) *	0.709649 (6.09E-2) *	0.214618 (8.11E-3) *	0.834999 (6.02E-3) *	<b>0.901471</b> (5.07E-2) *	0.784401 (6.12E-3) *
2	UF2	0.576268 (1.56E-2) *	0.673588 (9.42E-3) *	0.804848 (9.14E-3) *	0.870498 (1.07E-3) *	0.927186 (2.69E-3) *	0.911687 (1.94E-3) *	<b>0.932005</b> (1.41E-3) *
2	UF3	0.517992 (4.94E-3) *	0.536167 (1.09E-2) *	0.675722 (7.55E-3) *	0.628483 (2.77E-3) *	0.982468 (1.48E-3) *	0.980885 (3.36E-3) *	<b>0.989193</b> (1.14E-3) *
2	UF4	0.832407 (7.82E-3) *	0.904059 (4.44E-3) *	0.855195 (4.12E-3) *	0.890782 (1.98E-3) *	0.908135 (3.42E-2) *	0.906710 (5.15E-3) *	<b>0.915155</b> (1.53E-2) *
2	UF5	— (—) *	— (—) *	— (—) *	— (—) *	— (—) *	<b>0.199880</b> (9.99E-2) *	— (—) *
2	UF6	0.407692 (4.75E-2) *	0.706527 (4.21E-2) *	0.217237 (1.08E-1) *	— (—) *	0.503085 (2.10E-2) *	<b>0.701907</b> (6.42E-2) *	0.256224 (3.93E-2) *
2	UF7	0.606616 (1.86E-2) *	0.709033 (1.38E-2) *	0.725638 (1.48E-1) *	0.190892 (1.17E-2) *	0.842589 (7.20E-3) *	<b>0.898994</b> (3.27E-2) *	0.777471 (8.81E-3) *
2	ZDT1	0.854970 (8.69E-3) *	0.548507 (2.05E-2) *	0.568226 (4.09E-2) *	0.112146 (3.88E-2) *	0.998415 (1.55E-4) *	0.998462 (1.13E-4) *	<b>0.998609</b> (1.21E-4) *
2	ZDT2	0.758925 (2.05E-2) *	0.109237 (4.02E-2) *	0.110764 (5.31E-2) *	— (—) *	0.998239 (1.50E-4) *	0.998270 (1.49E-4) *	<b>0.998476</b> (1.74E-4) *
2	ZDT3	0.815271 (6.62E-2) *	0.512281 (1.82E-2) *	0.621983 (2.71E-2) *	0.217650 (3.62E-2) *	0.999277 (7.55E-5) *	0.999326 (5.44E-5) *	<b>0.999439</b> (5.62E-5) *
2	ZDT4	— (—) *	— (—) *	— (—) *	<b>0.998789</b> (4.13E-3) *	— (—) *	— (—) *	0.998615 (1.10E-4) *
2	ZDT6	— (—) *	— (—) *	— (—) *	— (—) *	0.998498 (1.69E-4) *	0.998639 (4.69E-5) *	<b>0.998766</b> (7.89E-5) *
3	DTLZ1	— (—) *	— (—) *	— (—) *	<b>0.587705</b> (1.48E-1) *	— (—) *	— (—) *	0.587705 (5.87E-1) *
3	DTLZ2	— (—) *	— (—) *	— (—) *	<b>0.981398</b> (2.04E-3) *	0.781343 (6.49E-2) *	0.976170 (3.24E-3) *	0.979655 (3.22E-3) *
3	DTLZ3	— (—) *	— (—) *	— (—) *	<b>0.390379</b> (—)	— (—) *	— (—) *	— (3.90E-1) *
3	DTLZ4	— (—) *	— (—) *	— (—) *	<b>1.004301</b> (1.61E-3) *	0.485969 (3.05E-1) *	0.976607 (1.14E-2) *	1.001749 (5.63E-3) *
3	DTLZ5	— (—) *	— (—) *	— (—) *	<b>0.998575</b> (1.93E-4) *	0.004036 (6.93E-2) *	0.982175 (5.72E-3) *	0.998159 (4.52E-4) *
3	DTLZ6	— (—) *	— (—) *	— (—) *	— (—) *	0.999158 (2.53E-4) *	0.999316 (1.58E-4) *	<b>0.999504</b> (1.14E-4) *
3	DTLZ7	0.279526 (2.50E-1) *	0.219555 (2.66E-2) *	0.477192 (2.50E-2) *	0.002625 (6.41E-3) *	<b>0.989964</b> (3.10E-3) *	0.989889 (2.55E-3) *	0.979407 (5.84E-3) *
3	WFG1	0.416198 (8.10E-3) *	0.570543 (3.66E-3) *	0.289553 (3.82E-3) *	0.579919 (4.74E-3) *	<b>0.604212</b> (1.12E-2) *	0.590271 (5.22E-3) *	0.602670 (7.99E-3) *
3	WFG2	0.552173 (5.18E-3) *	0.617007 (4.76E-3) *	0.655834 (5.87E-3) *	0.740452 (5.07E-3) *	0.868322 (1.10E-2) *	0.865291 (4.93E-2) *	<b>0.956430</b> (1.13E-2) *
3	WFG3	0.243353 (1.84E-2) *	0.476209 (9.00E-3) *	0.502668 (1.84E-2) *	0.564899 (1.61E-2) *	0.765131 (3.90E-2) *	0.804832 (2.64E-2) *	<b>0.948554</b> (1.29E-2) *
3	WFG4	0.357544 (7.68E-3) *	0.538518 (1.00E-2) *	0.509696 (1.05E-2) *	0.719935 (5.03E-3) *	0.806799 (2.02E-2) *	0.806814 (3.26E-2) *	<b>0.885714</b> (3.72E-2) *
3	WFG5	0.390448 (7.58E-3) *	0.601179 (1.16E-2) *	0.460440 (8.12E-3) *	0.678253 (5.92E-3) *	0.796938 (4.47E-2) *	0.793484 (3.09E-2) *	<b>0.835976</b> (2.78E-2) *
3	WFG6	0.316011 (1.03E-2) *	0.445367 (1.01E-2) *	0.451270 (1.29E-2) *	0.874272 (6.95E-3) *	0.845404 (1.03E-1) *	0.741775 (8.18E-2) *	<b>0.967580</b> (1.31E-2) *
3	WFG7	0.387126 (7.60E-3) *	0.567569 (8.38E-3) *	0.552353 (6.13E-3) *	0.662307 (1.42E-2) *	0.796639 (1.45E-2) *	0.800744 (1.63E-2) *	<b>0.856828</b> (2.75E-2) *
3	WFG8	0.327917 (6.15E-3) *	0.536088 (8.10E-3) *	0.453376 (7.62E-3) *	0.547306 (1.91E-2) *	0.739925 (2.35E-2) *	0.752790 (2.84E-2) *	<b>0.812080</b> (2.39E-2) *
3	WFG9	0.287493 (5.26E-3) *	0.369398 (1.82E-2) *	0.492735 (1.32E-2) *	0.824473 (2.25E-2) *	0.869764 (3.33E-2) *	0.841607 (2.40E-2) *	<b>0.903643</b> (3.07E-2) *
3	UF8	— (—) *	0.087400 (2.11E-2) *	0.296347 (5.62E-2) *	0.857926 (6.39E-4) *	0.855352 (1.53E-3) *	0.857292 (8.50E-4) *	<b>0.858015</b> (4.85E-4) *
3	UF9	0.002653 (8.61E-3) *	0.062455 (1.54E-2) *	0.353288 (5.55E-2) *	0.555565 (2.99E-3) *	<b>0.694577</b> (1.01E-2) *	0.642928 (6.99E-2) *	0.641168 (7.83E-3) *
3	UF10	— (—) *	— (—) *	— (—) *	0.854390 (3.84E-3) *	0.778501 (1.46E-1) *	0.690878 (1.10E-1) *	<b>0.855156</b> (2.40E-3) *

WOF-SMPSO and WFG-GDE3 is obtained). Additionally we want to point out the good performance on both, separable and nonseparable problems. Decomposition methods often face difficulties when the problems are nonseparable, since the interaction between the variables asks for a suitable grouping of the variables. The problems WFG2, WFG3, WFG6, WFG8, and WFG9 are nonseparable. Still we observe that WOF can achieve the same superior performance as on the separable problems, which let us assume that separability does not affect the WOF's performance much. It must be noted that the used ordered grouping mechanism is likely to separate the position- and distance-related variables into different groups. This lets WOF optimize them separately and results in a good performance of the nonseparable problems. However, further experiments using random grouping also showed a superior performance of the WOF algorithms on the separable and nonseparable WFG problems.

2) *ZDT*: In Table I, the best performance on the ZDT problems is obtained by WOF-algorithms. Only in the ZDT4, where all but the PSO-based algorithms fail to obtain any hypervolume value, the original SMPSO performs a little better than the WOF-SMPSO. This difference, however, might not

be of great importance, since both methods achieve a relative hypervolume of more than 99.8%.

3) *CEC 2009 Competition*: The results on the CEC2009 competition basically follow the same pattern. The WOF-SMPSO and the WOF-NSGA-II perform best and in almost all instances significantly better than the non-WOF algorithms. The only two exceptions are the UF8 and UF10 problems where the original SMPSO is on a par with its WOF version. That is, SMPSO is not significantly outperformed by its WOF version for UF8 and UF10.

4) *DTLZ*: On the DTLZ problems, the WOF versions of GDE3 and NSGA-II perform significantly better than their respective original parts. Also, all WOF algorithms perform better than the CCGDE3 algorithm. For most instances of DTLZ1–DTLZ5 in Table I, the best results are obtained by the SMPSO. On these problems, WOF does not further improve the performance of the SMPSO algorithm in the same way as it does of NSGA-II and GDE3, mainly due to a good performance of the original SMPSO in the first place. Although the results of the SMPSO are better than the WOF-SMPSO on these five problems, we also see that the results of WOF-SMPSO and SMPSO do not actually differ

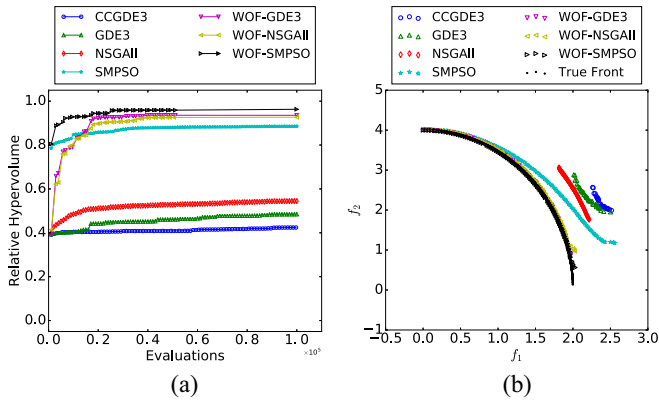


Fig. 7. Convergence and obtained solution sets on the WFG9 problem with  $n = 2000$  variables. (a) Convergence of rel. hypervolume. (b) Solution sets and Pareto-front.

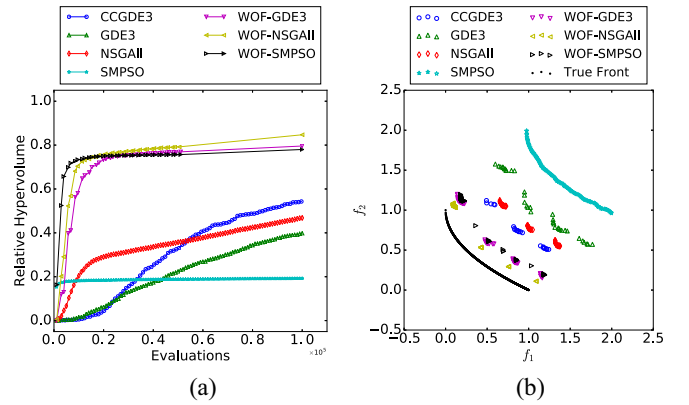


Fig. 8. Convergence and obtained solution sets on the CEC2009 UF1 problem with  $n = 2000$  variables. (a) Convergence of rel. hypervolume. (b) Solution sets and Pareto-front.

to a great extent, even though these differences are marked as statistically significant. The obtained relative hypervolumes range in the same order of magnitude, as both algorithms perform very similar to each other. This indicates that the SMPSO already performs well (to a certain degree) on the large-scale DTLZ1–DTLZ5 problems, thus the WOF version of it does not deliver the same amount of improvement as for the other algorithms. At the same time, we can observe that the performance of the NSGA-II and GDE3 methods are improved significantly, as they perform poor without WOF, but much better when WOF is used. For the DTLZ6 and DTLZ7 problems, we observe the same superior performance of all WOF algorithms as for the other problem families in Table I. We can conclude for the DTLZ problems that the WOF does improve the performance of previously bad-performing algorithms (NSGA-II and GDE3) in the same way as in the case of the UF and WFG benchmarks, and does on the other hand deliver comparable results to the original algorithm (SMPSO), if this one does already deal well with the large-scale problem.

A special consideration goes to the CCGDE3, since it is an algorithm that was specifically designed to work well on large-scale problems. In the experiments performed in this section, it could not perform of the same level with any of the WOF algorithms on any on the test problems. The comparison using the original settings of the CCGDE3 in Section VI-D will support this in further detail.

Concerning the diversity of solutions, Figs. 7–12 and also Figs. 14–41 of the supplementary material illustrate that the obtained solutions are well-spread for several test problems. Nevertheless, the final populations obtained by our approach do not always cover the entire Pareto-fronts. On the other hand, we observe much better convergence performance compared to the other algorithms. This observation clearly shows the difficulty to find a balance between convergence and diversity for large-scale MOPs. Our approach seems to be somewhat convergence-oriented. Diversity improvement is an important future research topic including the use of uniformly specified reference lines instead of the crowding distance for selecting solutions (see Section IV-C).

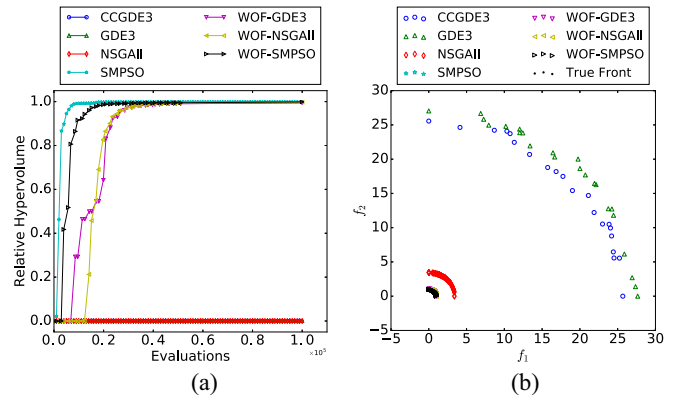


Fig. 9. Convergence and obtained solution sets on the DTLZ2 problem with  $n = 1000$  variables. (a) Convergence of rel. hypervolume. (b) Solution sets and Pareto-front.

### C. Scaling the Number of Variables

Table I gave some detailed results for  $n = 1000$  variables. In addition to the analysis that we gave for each of the test problems until now, we examined how these results change with the dimension of the search space. For each setting of  $n \in \{40, 80, 200, 600, 1000, 2000\}$ , we perform the same experiment as in the previous section. Due to limited space, Tables 15 and 16 of the supplementary material to this paper give an overview over these experiments. In these tables, for every combination of problem and number of variables we list the algorithm that achieved the highest as well as second highest median hypervolume values. An asterisk is used to indicate whether the respective values are significantly better than the next-best performance.

As expected, for small numbers of variables, the original optimization methods SMPSO, NSGA-II, and GDE3 work best on most problems, as seen in Tables 15 and 16 of the supplementary material in the experiments with  $n = 40$  and  $n = 80$ . The exception is the WFG problem family, where the WOF versions already work better than the original algorithms for these relatively small problems instances, although the differences are not always statistically significant in these cases.



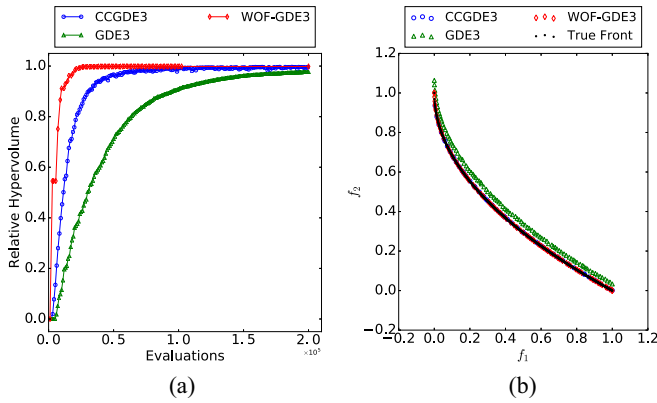


Fig. 10. Comparison with CCGDE3: convergence and obtained solution sets on the ZDT1 problem with  $n = 500$  variables. (a) Convergence of rel. hypervolume. (b) Solution sets and Pareto-front.

When the number of variables is increased, we observe a superior performance of the WOF-methods, especially the WOF-SMPSO. In the two-objective experiments (Table 15 of the supplementary material), for  $n = 600$ ,  $n = 1000$  (which corresponds to the results in Table I), and  $n = 2000$ , the WOF algorithms are placed best for all WFG, ZDT, and UF problems, with the exception of ZDT4.

In Figs. 7–9, we show the convergence rate and obtained solution sets on three problem instances. The runs shown are the ones that achieved the median hypervolume of each respective algorithm among all 51 runs. Fig. 7 shows the WFG9 problem with 2000 variables. We observe not only a superior performance in terms of closeness to the Pareto-front as shown in the previous tables, but also a very fast convergence rate especially in the beginning of the optimization. The same can be seen in Fig. 8 for the UF1 problem with 2000 variables. All three WOF algorithms obtain a much closer approximation of the Pareto-front. Additionally the WOF algorithms only need a very small share of the total function evaluation to reach this approximation. Finally, in Fig. 9 we observe the DTLZ2 performance for  $n = 1000$  variables. As seen before, the WOF does not perform in the same superior way when used with the SMPSO algorithm on the DTLZ problems. However, the convergence as seen in Fig. 9(a) shows that all three WOF algorithms do still outperform all but the SMPSO algorithm in terms of solution quality as well as convergence rate.

In conclusion, we observe that for any LSO problem, the WOF methods can outperform the traditional GDE3 and NSGA-II methods as well as the large-scale method CCGDE3 significantly. The performance of the SMPSO lies on par with the WOF-SMPSO for some of the DTLZ problems, but the WOF method also increases its performance significantly on the other benchmark families. Overall, the results of the WOF do not only show a superior solution quality, but also a very fast convergence rate.

#### D. Comparison With CCGDE3

As described above, CC has been used frequently for LSO, especially for single-objective problems. A few works also

TABLE II  
MEDIAN RELATIVE HYPERVOLUME VALUES AND IQRs OF THE ZDT PROBLEMS WITH DIFFERENT NUMBERS OF VARIABLES. SETTINGS USED ARE THE SAME AS IN [11]. BEST PERFORMANCE IS SHOWN IN BOLD FONT. AN ASTERISK INDICATES STATISTICAL SIGNIFICANCE COMPARED TO THE RESPECTIVE BEST METHOD

	CCGDE3	GDE3	WOF-GDE3
<b>n = 200</b>			
ZDT1	0.996298 (3.91E-3) *	0.998319 (6.82E-5)	<b>0.998346</b> (2.13E-4)
ZDT2	0.983671 (9.13E-3) *	0.997598 (1.88E-4) *	<b>0.998198</b> (3.25E-4)
ZDT3	0.958335 (6.09E-2) *	0.999220 (3.31E-5)	<b>0.999239</b> (1.18E-4)
ZDT4	— (—) *	— (—) *	<b>0.545522</b> (—)
ZDT6	0.982064 (8.55E-3) *	0.858491 (6.72E-3) *	<b>0.998698</b> (2.58E-4)
<b>n = 500</b>			
ZDT1	0.995791 (3.54E-3) *	0.976890 (1.99E-3) *	<b>0.998392</b> (1.53E-4)
ZDT2	0.985697 (1.01E-2) *	0.960381 (2.26E-3) *	<b>0.998217</b> (2.03E-4)
ZDT3	0.929806 (6.71E-2) *	0.951782 (4.13E-3) *	<b>0.999252</b> (1.04E-4)
ZDT4	— (—) *	— (—) *	<b>0.545522</b> (—)
ZDT6	0.862895 (1.13E-2) *	— (—) *	<b>0.998573</b> (3.30E-4)
<b>n = 1000</b>			
ZDT1	0.984426 (4.46E-3) *	0.783685 (1.90E-2) *	<b>0.998382</b> (2.19E-4)
ZDT2	0.966051 (1.46E-2) *	0.549738 (3.50E-2) *	<b>0.998204</b> (3.98E-1)
ZDT3	0.922063 (6.41E-2) *	0.696589 (1.63E-2) *	<b>0.999267</b> (1.54E-4)
ZDT4	— (—) *	— (—) *	<b>0.545522</b> (—)
ZDT6	0.453370 (5.22E-2) *	— (—) *	<b>0.998545</b> (2.91E-4)
<b>n = 2000</b>			
ZDT1	0.932251 (6.91E-3) *	0.371685 (2.40E-2) *	<b>0.998410</b> (2.05E-4)
ZDT2	0.892224 (1.77E-2) *	— (—) *	<b>0.998161</b> (3.98E-1)
ZDT3	0.761271 (2.04E-1) *	0.384368 (1.15E-2) *	<b>0.999264</b> (8.57E-5)
ZDT4	— (—) *	— (—) *	<b>0.545522</b> (—)
ZDT6	— (—) *	— (—) *	<b>0.998367</b> (3.86E-4)
<b>n = 3000</b>			
ZDT1	0.838997 (9.71E-3) *	0.193257 (2.31E-2) *	<b>0.998411</b> (1.58E-4)
ZDT2	0.774619 (3.05E-2) *	— (—) *	<b>0.998220</b> (3.98E-1)
ZDT3	0.575661 (1.58E-1) *	0.262717 (1.33E-2) *	<b>0.999242</b> (1.31E-4)
ZDT4	— (—) *	— (—) *	<b>0.545522</b> (—)
ZDT6	— (—) *	— (—) *	<b>0.998292</b> (6.89E-4)
<b>n = 4000</b>			
ZDT1	0.720947 (1.54E-2) *	0.104674 (1.66E-2) *	<b>0.998407</b> (2.20E-4)
ZDT2	0.633076 (2.28E-2) *	— (—) *	<b>0.998246</b> (3.98E-1)
ZDT3	0.472919 (1.52E-1) *	0.199042 (9.78E-3) *	<b>0.999227</b> (1.16E-4)
ZDT4	— (—) *	— (—) *	<b>0.545522</b> (—)
ZDT6	— (—) *	— (—) *	<b>0.998138</b> (6.87E-4)
<b>n = 5000</b>			
ZDT1	0.614349 (2.17E-2) *	0.047964 (1.06E-2) *	<b>0.998428</b> (1.44E-4)
ZDT2	0.486880 (2.45E-2) *	— (—) *	<b>0.998159</b> (3.98E-1)
ZDT3	0.384726 (1.61E-1) *	0.160194 (8.56E-3) *	<b>0.999234</b> (1.18E-4)
ZDT4	— (—) *	— (—) *	<b>0.545522</b> (—)
ZDT6	— (—) *	— (—) *	<b>0.998120</b> (1.00E-3)

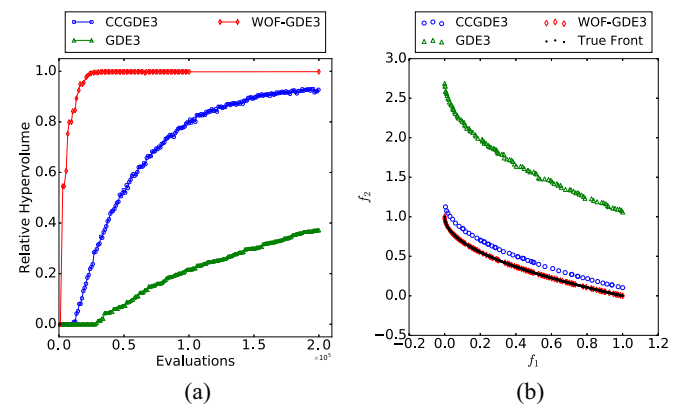


Fig. 11. Comparison with CCGDE3: convergence and obtained solution sets on the ZDT1 problem with  $n = 2000$  variables. (a) Convergence of rel. hypervolume. (b) Solution sets and Pareto-front.

used CC for multiobjective optimization and performed experiments using the ZDT [5] suite. One of the most recent advances is the CCGDE3 algorithm presented in [11], where the CC concept is applied to the GDE3 algorithm.



In the previous subsection, we observe in Table 15 of the supplementary material that for only one small-scale problem instance of the UF5 problem with  $n = 80$  variables and two objectives CCGDE3 could win against the other methods. However, except for this 80-variable UF5 problem, CCGDE3 is not performing best for any test problem in Table I and Tables 15 and 16 of the supplementary material.

In this section we will further compare the performance of the WOF with CCGDE3. As experimental benchmark, the ZDT problems 1–4 and 6 were used with the numbers of decision variables ranging from 200 to 5000. To compare the WOF with the CCGDE3 method, we perform all experiments using the same configurations and parameter settings as in the original work [11]. We compare the performance of the original GDE3 algorithm, CCGDE3, and WOF-GDE3 using the same test problems (ZDT1–3 and 6) that were used in [11], as well as the ZDT4 problem.

The parameter settings differ from Section VI-A in the following. The number of groups  $\gamma$  is set to 2 in CC and WOF. The grouping mechanism used for both is a random grouping of the variables. Furthermore, since the optimal solutions of the ZDT problems lie on the boundaries of the variables domains, for fair comparison we use the transformation function  $\psi_3$  as described in Section V-B. In this way, the need for a repair mechanism is almost eliminated. When infeasible values are created using the obtained weights on other solutions in the population, the respective variable values are set to random values instead of the extremal ones.

The results of this experiment can be seen in Table II. Even for rather small problem instances ( $n = 200$ ) the WOF-GDE3 can outperform the two other methods. Only for the ZDT1 and ZDT2 problems, the original GDE3 algorithm does not perform significantly worse.

With increasing the number of variables we see that the original GDE3 cannot find a good approximation of the Pareto-front for any problem. The CCGDE3 starts to outperform the original GDE3 when the number of variables grows larger. However, for all problem instances with  $n \geq 500$ , the WOF algorithm clearly obtains the highest median hypervolume in all test problems with statistical significance (with the one exception of ZDT2 with 4000 variables). This shows that for a large number of decision variables the WOF algorithm is clearly superior to the CC framework on the ZDT benchmarks. This adds to the results seen in Sections VI-B and VI-C, where we saw that the superiority of WOF is already present for much smaller values of  $n$  when other problems than the ZDT are used.

Another aspect that is notable is once again the convergence rate of the WOF algorithm. The difference in performance for the ZDT1 problem is shown in Figs. 10–12. We plotted for each algorithm the respective run that achieved the median hypervolume of all 51 runs for  $n = 500$ ,  $n = 2000$ , and  $n = 5000$  decision variables. Additionally, the convergence rate is plotted next to it. In these figures, one can clearly see that the WOF algorithm was able to obtain an almost perfect approximation of the Pareto-front with a good diversity in all cases. In contrast, the original GDE3 and the CCGDE3 are not able to obtain any good approximation of the Pareto-optimal

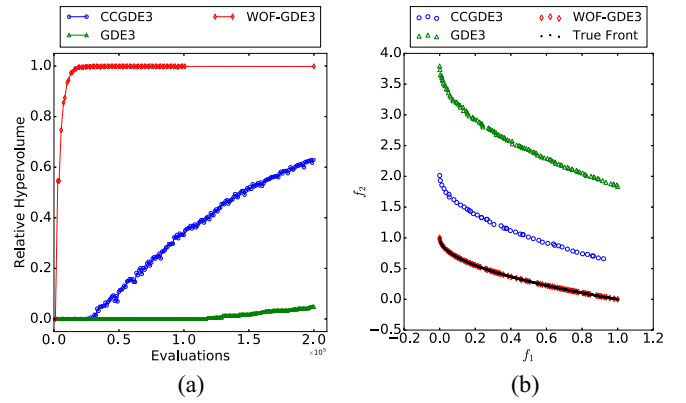


Fig. 12. Comparison with CCGDE3: convergence and obtained solution sets on the ZDT1 problem with  $n = 5000$  variables. (a) Convergence of rel. hypervolume. (b) Solution sets and Pareto-front.

TABLE III  
COMPARISON OF GROUPING SCHEMES AND TRANSFORMATION FUNCTIONS. SHOWN ARE THE NUMBERS OF TIMES THE RESPECTIVE COMBINATION ACHIEVES THE BEST/SECOND BEST/ WORST MEDIAN REL. HYPERVOLUME OVER ALL TESTED PROBLEMS

		Transformation function		
		$\psi_1$	$\psi_2$	$\psi_3$
Grouping	Random	1 / 1 / 8	3 / 1 / 2	1 / 1 / 7
	Ordered	4 / 4 / 3	3 / 6 / 0	2 / 4 / 4
	Linear	4 / 2 / 2	2 / 2 / 2	2 / 4 / 3
	Differential	4 / 1 / 3	7 / 0 / 1	2 / 5 / 1

solutions for  $n > 500$ . In all cases, we also see that the convergence rate is extremely high for the WOF-GDE3. In all three plots, it reaches a relative hypervolume of over 99% already after less than 12% of the total function evaluations, no matter how many variables are used.

A special consideration goes to the ZDT4 problem. It is the only tested problem that allows negative variable values, which limits the reachable search space of the WOF algorithm. Since we used the  $\psi_3$  transformation function, as explained before the weights can only map positive to positive and negative to negative variable values. As a result, a part of the search space might not be reachable in the weight optimization steps of WOF. Nevertheless, the performance of the WOF on the ZDT4 problem is significantly higher than that of the CCGDE3 for any tested number of variables. While the CCGDE3 does not obtain any hypervolume value with the used reference point, the WOF always obtains the same hypervolume value (in all runs, therefore the IQR is nonexistent). A closer look in the results showed that in fact the WOF-GDE3 always finds exactly one (same) solution that is part of the true Pareto-front in every run. Therefore, its performance can be regarded as extremely reliable for ZDT4, while in contrast no diversity exists. This, however, is not a direct weakness of the WOF-method, because we saw that the WOF-SMPSO performed well on the ZDT4 problem before (see Table I).

### E. Comparison With MOEA/DVA

In this part, we compare the performance of our WOF method with MOEA/DVA [12].<sup>2</sup> This approach focusses

<sup>2</sup>Codes are obtained from the original authors' webpage: <http://web.xidian.edu.cn/home/fliu/lunwen.html>.

on finding optimal groupings of the variables by dividing them into diversity-related and convergence-related variables. In comparison with our proposed method, some interesting results can be observed. The tables and figures supporting these can be found in the supplementary material to this paper. We compared the performance of the best found WOF-version from the previous experiments, the WOF-SMPSO, with MOEA/DVA on the UF1–UF10 as well as the two-objective WFG1–WFG5 and WFG7 and the three-objective WFG1–WFG5 problems using 1000 decision variables in all instances.

The experiments in this section differ from the previous ones in only one aspect which is the number of function evaluations used. The MOEA/DVA uses the majority of its total functions evaluations in the preliminary variable grouping method, before the actual optimization starts. In our experiments, the grouping mechanism of MOEA/DVA for 1000-variable problems consumed around 9 000 000 evaluations. Given that all previous experiments with WOF in this paper only use 100 000 evaluations in total, the comparison might not be fair in this case. For this reason we decided to provide all algorithms a total of 10 000 000 evaluations for the experiments in this section. To compare, the original article of MOEA/DVA [12] used between 1 200 000 and 3 000 000 evaluations in their experiments with the 200-variable problems. The parameters of the MOEA/DVA were the same as found in the original MOEA/DVA paper with the exception of adjusting the population sizes to the same values as the ones of SMPSO and WOF-SMPSO to ensure a fair comparison using the hypervolume indicator. Furthermore, the distribution indexes of crossover and mutation are set to the same values as described in Section VI-A. No other changes were made to the original sourcecode provided by Ma *et al.* [12].

The obtained median hypervolume and IQR values are shown in Tables 13 and 14 in the supplementary material of this paper, and the solution sets and convergence plots for the two-objective problems can be found in Figs. 1–13 in the supplementary material. The results show that on the (two-objective) UF1–UF7 and the UF9 problems, the MOEA/DVA performs better than the WOF-SMPSO. On the other hand, the WOF-SMPSO significantly outperforms MOEA/DVA on all tested WFG problem instances with two and three objectives and the UF10 problem. Partly this difference results from a slightly worse performance of the WOF algorithm on the UF problems; however, we also observed a large decrease in the performance of MOEA/DVA on the WFG problems in comparison to UF.

A second observation regards the convergence rate. In contrast to the MOEA/DVA, which uses 90% of the total evaluations for an initial grouping phase, the WOF-SMPSO does immediately start with the optimization process and can therefore deliver solution sets of reasonable quality after just a fraction of the total evaluations. In all used WFG2, WFG4, WFG5, and WFG7 as well as UF3 instances the WOF-SMPSO provides a hypervolume value of over 0.95 after just 10% (1 000 000) of the total function evaluations, and in the WFG1 and WFG3 it obtains a hypervolume very close to the final value after this amount. For all other problems we

observe a more gradual convergence, however, the advantage remains that even with a very small amount of evaluations, we can already obtain results of certain quality using the WOF-SMPSO (as also seen in Section VI-B where we used only 100 000 evaluations). In contrast, while the MOEA/DVA outperforms WOF on most of the UF problems, these results can only be obtained after a large computational effort, since no solution sets are produced by MOEA/DVA during its grouping phase. As a conclusion, there are some problems where the MOEA/DVA outperforms WOF-SMPSO, but with the disadvantage of using a much larger computational budget, while on some other problems the WOF-SMPSO performs significantly better with a much smaller computation effort.

#### F. Influence of Grouping and Transformation Methods

In this section we will examine the effects of different grouping mechanisms and transformation functions. We implement the four grouping mechanisms and the three transformation functions explained in Section V. The WOF-SMPSO algorithm is used for comparison. For each combination of a grouping mechanism and a transformation function, we perform the same experiment as in Section VI-B on the WFG1–WFG4, DTLZ1–DTLZ4, and ZDT problems with two and three objectives. As we are only interested in the efficiency of the mechanisms, the DG was precomputed for each problem using an  $\epsilon$ -value of  $10^{-10}$ , i.e., the consumed evaluations were not counted to those of the optimization process.

Table III lists the number of problems (out of the 21 test problems) for which the respective combination performed best, second best and worst. We observe that the algorithms using the  $p$ -value transformation ( $\psi_2$ ) and the multiplication transformation  $\psi_1$  perform on average better than the  $\psi_3$  transformation function. Comparing the grouping mechanisms, we see that the random grouping seems to perform worse than the other mechanisms. The best combination was obtained when using the DG method together with the  $p$ -value ( $\psi_2$ ) transformation.

We take a closer look at the performance of DG. As a single-objective grouping strategy, it is based on only the first objective in this paper. Therefore, we did not expect its high performance for the MOPs. If we look closer for instance at the ZDT problems, which are all separable, we observe that all their variables are in the same (single) group (since all separable variables by default are grouped together by the original DG algorithm). This results in the optimization of just one weight value for all variables. For all  $i = 2, \dots, n$ , reducing the value of a variable  $x_i$  in a ZDT problem is improving solution quality in the second objective component. Therefore, we can expect a speed-up in convergence by using just one single weight.

The better performance of the linear grouping compared to the random one (in  $\psi_1$  and  $\psi_3$ ) can be explained by taking a look at the variables of the WFG problems. Since out of the 1000 variables the first quarter of them are position-related variables, a linear grouping with  $\gamma = 4$  groups results in a splitting of position and distance variables. This might be a benefit compared to the random grouping and therefore leads to a better overall performance. Our observations further show

that DG, although just regarding one objective, can approximately divide position and distance variables of some of the WFG problems into different groups.

Overall, no single combination is clearly superior in the majority of the 21 test problems used in this experiment. These observations show that the choice of a good grouping mechanism can strongly affect the performance of the WOF method. Finding a single mechanism that fits all problems might be a difficult task. It can be expected, however, that by developing an advanced multiobjective grouping mechanism, the performance can be further improved.

### G. Sensitivity Analysis

In this part, we take a closer look at the different parameters of the WOF and analyze the sensitivity to certain changes in these parameter values. The tested parameters are: 1)  $q$ , the number of chosen solutions  $\vec{x}_k$ ; 2)  $\gamma$ , the number of groups used in the ordered grouping mechanism; 3)  $\delta$ , the share of the total function evaluations used for the weight optimization; 4)  $p$ , the value used in the  $\psi_2$  transformation function; 5)  $t_1$ , the amount of evaluations used for each optimization of the original problem (line 4 in Algorithm 1); and 6)  $t_2$ , the amount of evaluations used for each optimization of the transformed problem (line 5 in Algorithm 2). For each parameter we tested five different values including values higher and lower than those described previously in Section VI-A. All other parameters remain unchanged. The sensitivity experiments are done with the WOF-SMPSO and the WOF-NSGA-II algorithms using the UF, WFG, DTLZ, and ZDT benchmark functions with two and three objectives, where applicable, and 21 independent runs are performed for each configuration. The detailed results can be found in Tables 1–12 of the supplementary material of this paper.

For the parameters  $t_1$  and  $t_2$  we observe a low sensitivity in general. While the performance varies when ranging  $t_1$  from 400 to 2000 and  $t_2$  from 100 to 900, the differences in the hypervolume are in most cases not statistically significant for both tested algorithms. For the value of  $p$  in the transformation function, in most problem instances a significantly lower performance was observed for a very small value of 0.05 in both algorithms. For the WOF-SMPSO, performance remained stable for values between 0.2 and 0.5, while the NSGA-II achieved on average better results with higher values of  $p$ . The value of  $\delta$  was altered between 0.1 and 1.0, and we observed an average increase in performance for higher values of  $\delta$ , especially when the NSGA-II is used within our framework. On the other hand the results when using the SMPSO also show, that the optimal value for  $\delta$  might depend on the used optimizer in WOF as well as the problem to be optimized. Future work on the framework should therefore include an automatic detection of the starting point for the traditional optimization phase. For the parameter  $\gamma$ , the number of groups, we observe that the WOF-SMPSO and WOF-NSGA-II perform significantly worse if the values of  $\gamma$  are too high ( $\gamma = 50$ ) or too low ( $\gamma = 2$ ). These observations, however, were made for the described  $p$ -value transformation ( $\psi_2$ ), which we do not claim to be a perfect grouping mechanism. A more suitable grouping mechanism (e.g., from [12] or [29]) could be used

within WOF to obtain more reliable results, since the optimal number of groups might depend on each specific problem. Lastly, for the parameter  $q$ , the number of chosen solutions, differences in the performance were observed by varying  $q$ , but most of these were not significant with the exception of choosing extreme values ( $q = 1$  or  $q = 8$ ). This result was to be expected as a different number of  $q$  can be used to balance between convergence and diversity.

## VII. CONCLUSION

In this paper, we explained and examined the idea proposed in [3] in detail. We introduce the use of a framework for multiobjective optimization that is able to improve the performance of multiobjective optimization algorithms for many-variable problems. The proposed WOF is not making use of any coevolution as many other approaches in this area and is designed to employ another multiobjective optimization algorithm while transforming the problem during the optimization. We explained the underlying problem transformation mechanism and the algorithm in detail and examined its implications for the optimization process. Different grouping and transformation functions were introduced. Through experimental evaluation we showed significant improvements of the performance on various benchmark problems compared to classical optimization methods as well as existing large-scale approaches. The best performance was achieved when the SMPSO algorithm was used in the proposed framework. Overall, the WOF showed significant improvements in solution quality and convergence rate for almost all tested benchmark problems.

Future work might involve the implementation of better strategies for choosing the used solutions  $\vec{x}_k$  for problem transformations. An important issue in future research might also be the development of more advanced multiobjective grouping mechanisms, since current intelligent grouping methods are mostly single-objective oriented. Furthermore, since the  $q$  different transformed problems are optimized independently, this paper can easily be extended to a parallel version in the future.

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