

Particle Swarms with Dynamic Topologies and Conservation of Function Evaluations

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Abstract: This paper proposes a general framework for structuring dynamic Particle Swarm populations and uses a conservation of function evaluations strategy to increase the convergence speed. The population structure is constructed by placing the particles on a 2-dimensional grid of nodes, where they interact and move according to simple rules. During the running time of the algorithm, the von Neumann neighborhood is used to decide which particles influence each other when updating their velocity and position. Each particle is updated in each time-step but they are evaluated only if there are other particles in their neighborhood. A set of experiments demonstrates that the dynamics imposed by the structure provides a more consistent and stable behavior throughout the test set when compared to standard topologies, while the conservation of evaluations significantly reduces the convergence speed of the algorithm. Furthermore, the working mechanisms of the proposed structure are very simple and, except for the size of the grid, they do not require parameters and tuning.

1 INTRODUCTION

Kennedy and Eberhart (1995) proposed the Particle Swarm Optimization (PSO) algorithm for binary and real-valued function optimization, a method inspired by the swarming and social behavior of bird flocks and fish schools. Since then, PSO has been applied with success to a wide range of problems but the proper balance between exploration (global search) and exploitation (local search) is still an open problem that motives several lines research on the various mechanisms that control the algorithm's performance.

Population topology is one of PSO's components that affect the balance between exploration and exploitation and the convergence speed and accuracy of the algorithm. In the context of particle swarms, topology is the structure that defines the connections between the particles and consequently the flow of information through the population. The

reason why particles are interconnected is the core of the algorithm: the particles communicate so that they acquire information on the regions explored by other particles. In fact, it has been claimed that the uniqueness of the PSO algorithm lies in the interactions of the particles (Kennedy and Mendes, 2002). The population can be structured on any possible topology, from sparse to dense (or even fully connected) graphs, with different degrees of connectivity and clustering. The classical and most used population structures are the *lbest* (which connects the individuals to a local neighborhood) and the *gbest* (in which each particle is connected to every other individual). These topologies are well-studied and the major conclusions are that *gbest* is fast but is frequently trapped in local optima, while *lbest* is slower but converges more often to the neighborhood of the global optima.

Since the first experiments on *lbest* and *gbest* structures, researchers have tried to design networks that hold the best traits given by each structure

(Parsopoulos and Vrahatis, 2004). Some studies also try to understand what makes a good structure: for instance, Kennedy and Mendes (2002) investigate several types of topologies and recommend the use of a lattice with von Neumann neighborhood (which results in a connectivity degree between that of *lbest* and *gbest*).

Recently, dynamic structures have been tested in order to improve the algorithm's adaptability to different fitness landscapes and overcome the rigidity of static structures, like in (Liang et al., 2006). Fernandes et al. (2003) try a different approach and propose a dynamic and partially connected von Neumann structure with Brownian motion. This paper uses this model but introduces a strategy for the conservation of function evaluations (Majercik, 2013) with the aim of taking advantage of the underlying structure and reduce convergence speed. Furthermore, a formal description of the dynamic network is given here, opening the way for more sophisticated dynamics.

In the proposed topology, n particles are placed in a 2-dimensional m -nodes grid where $m > n$. Every time-step, each individual checks its von Neumann neighborhood and, as in the standard PSO, updates its velocity and position using the information given by the neighbors. However, while the connectivity degree (number of numbers, considering the particle itself) of the von Neumann topology is $k = 5$, the degree of the proposed topology is variable: $k \leq 5$. Furthermore, the structure is dynamic: in each time-step, every particle updates its position on the grid (which is a different concept from the position of the particle on the fitness landscape) according to a pre-defined rule that selects the destination node. The movement rule, which is implemented locally and without any knowledge on the global state of the system, can be based on stigmergy (Grassé, 1959) or Brownian motion.

As stated above, the connectivity degree k of each particle in each time-step is variable and lies in the range $1 \leq k \leq 5$. Depending on the size of the grid, there are, in each time-step, a number of particles with $k = 1$. These particles without neighbors (except the particle itself) do not learn from any local neighborhood at that specific iteration. Therefore, it is expected that they continue to follow their previous trajectory in the fitness landscape. Taking into account these premises, the algorithm proposed in this study does evaluate the position of the particles when $k = 1$. Regardless of the loss of informant intrinsic to a *conservation of evaluations* policy, we hypothesize that the strategy is

particularly suited for the proposed dynamic topology (in which the particles are sometimes isolated from the flow of information) and the number of function evaluations required for meeting the stop criteria can be significantly reduced. Furthermore, it is the structure of the population and the position of the particles at a specific time-step that decides the application of the conservation rule and not any extra parameter or pre-defined decision rule.

A classical PSO experimental setup is used for the tests and the results demonstrate that the proposed algorithm consistently improves the speed of convergence of the standard von Neumann structure without degrading the quality of solutions. The experiments also demonstrate that the introduction of the conservation strategy reduces significantly the convergence speed without affecting the quality of the final solutions.

The remaining of the paper is organized as follows. Section 2 describes PSO and gives an overview on population structures for PSOs. Section 3 gives a formal description of the proposed structure. Section 4 describes the experiments and discusses the results and, finally, Section 5 concludes the paper and outlines future research.

2 BACKGROUND REVIEW

PSO is described by a simple set of equations that define the velocity and position of each particle. The position of the i -th particle is given by $\vec{X}_i = (x_{i,1}, x_{i,2}, \dots, x_{i,D})$, where D is the dimension of the search space. The velocity is given by $\vec{V}_i = (v_{i,1}, v_{i,2}, \dots, v_{i,D})$. The particles are evaluated with a fitness function $f(\vec{X}_i)$ and then their positions and velocities are updated by:

$$v_{i,d}(t) = v_{i,d}(t-1) + c_1 r_1 (p_{i,d} - x_{i,d}(t-1)) + c_2 r_2 (p_{g,d} - x_{i,d}(t-1)) \quad (1)$$

$$x_{i,d}(t) = x_{i,d}(t-1) + v_{i,d}(t) \quad (2)$$

where p_i is the best solution found so far by particle i and p_g is the best solution found so far by the neighborhood. Parameters r_1 and r_2 are random numbers uniformly distributed in the range $[0,1]$ and c_1 and c_2 are acceleration coefficients that tune the relative influence of each term of the formula. The first term is known as the *cognitive part*, since it relies on the particle's own experience. The last term

is the social part, since it describes the influence of the community in the trajectory of the particle.

In order to prevent particles from stepping out of the limits of the search space, the positions $x_{i,d}(t)$ are limited by constants that, in general, correspond to the domain of the problem: $x_{i,d}(t) \in [-Xmax, Xmax]$. Velocity may also be limited within a range in order to prevent the *explosion* of the velocity vector: $v_{i,d}(t) \in [-Vmax, Vmax]$.

For achieving a better balancing between local and global search, Shi and Eberhart (1998) added the inertia weight ω as a multiplying factor of the first term of equation 1. This paper uses PSOs with inertia weight.

The neighborhood of the particle defines the value of p_g and is a key factor in the performance of PSO. Most of the PSOs use one of two simple sociometric principles for defining the neighborhood network. One connects all the members of the swarm to one another, and it is called *gbest*, where *g* stands for *global*. The degree of connectivity of *gbest* is $k = n$, where n is the number of particles. Since all the particles are connected to every other and information spreads easily through the network, the *gbest* topology is known to converge fast but unreliably (it often converges to local optima).

The other standard configuration, called *lbest* (where *l* stands for *local*), creates a neighborhood that comprises the particle itself and its k nearest neighbors. The most common *lbest* topology is the ring structure, in which the particles are arranged in a ring structure (resulting in a degree of connectivity $k = 3$, including the particle). The *lbest* converges slower than the *gbest* structure because information spreads slower through the network but for the same reason it is less prone to converge prematurely to local optima. In-between the ring structure with $k = 3$ and the *gbest* with $k = n$ there are several types of structure, each one with its advantages on a certain type of fitness landscapes. Choosing a proper structure depends on the target problem and also on the objectives or tolerance of the optimization process.

Kennedy and Mendes (2002) published an exhaustive study on population structures for PSOs. They tested several types of structures, including the *lbest*, *gbest* and von Neumann configuration with radius 1 (also known as *L5* neighborhood). They also tested populations arranged in randomly generated graphs. The authors conclude that when the configurations are ranked by the performance the

structures with $k = 5$ (like the *L5*) perform better, but when ranked according to the number of iterations needed to meet the criteria, configurations with higher degree of connectivity perform better. These results are consistent with the premise that low connectivity favors robustness, while higher connectivity favors convergence speed (at the expense of reliability). Amongst the large set of graphs tested in (Kennedy and Mendes, 2002), the Von Neumann with radius 1 configuration performed more consistently and the authors recommend its use.

Alternative topologies that combine standard structures' characteristics or introduce some kind of dynamics in the connections have been also proposed. Parsopoulos and Vrahatis (2004) describe the unified PSO (UPSO), which combines the *gbest* and *lbest* configurations. Equation 1 is modified in order to include a term with p_g and a term with p_i and a parameter balances the weight of each term. The authors argue that the proposed scheme exploits the good properties of *gbest* and *lbest*. Peram *et al.* (2003) proposed the fitness-distance-ratio-based PSO (FDR-PSO), which defines the "neighborhood" of a particle as its k closest particles in the population (measured by the Euclidean distance). A selective scheme is also included: the particle selects nearby particles that have also visited a position of higher fitness. The algorithm is compared to a standard PSO and the authors claim that FDR-PSO performs better on several test functions. However, the FDR-PSO is compared only to a *gbest* configuration, which is known to converge frequently to local optima in the majority of the functions of the test set. More recently, a comprehensive-learning PSO (CLPSO) was proposed (Liang *et al.* 2006). Its learning strategy abandons the global best information and introduces a complex and dynamic scheme that uses all other particles' past best information. CLPSO can significantly improve the performance of the original PSO on multimodal problems. Finally, Hsieh *et al.* (2009) use a PSO with varying swarm size and solution-sharing that, like in (Liang *et al.* 2006), uses the past best information from every particle.

A different approach is given in (Fernandes *et al.*, 2013). The authors describe a structure that is based on a grid of m nodes (with $m > n$) on which the particles move and interact. The von Neumann neighborhood is checked for choosing the informants of the cognitive part of equation 1. Since $m > n$, the number of neighbors lies in the range $1 \leq k \leq 5$ and

the distribution of the particles on the grid at a given time-step defines the PSO structure at that precise iteration. The results demonstrate that the proposed structure performs consistently throughout the test set, improving the performance of other topologies in the majority of the scenarios and under different performance evaluation criteria. Furthermore, the structure is very simple and only the grid size needs to be set before the run (the authors suggest 1:2 ratio between the swarm size and the number of nodes). However, the structure itself and the distribution of the particles on the grid suggest that there is still room for improvement, namely of convergence speed, which is a critical aspect when optimizing real-world functions.

In the proposed structure, a particle, at a given time-step, may have no neighbors except itself. The isolated particles will continue to follow its previous trajectory, based on their current information, until they find another particle in the neighborhood. Therefore, we intend to investigate if the loss of information caused by not evaluating these particles is overcome by the payoff in the convergence speed.

Common ways of addressing the computational cost of evaluating solutions in hard real-world problems are function approximation (Landa-Becerra et al., 2008), fitness inheritance (Reyes-Sierra and Coello Coello, 2007) and conservation of evaluations (Majercik, 2013). Due to the underlying structure of the proposed algorithm, we have tested a conservation policy similar to the GREEN-PSO proposed by Majercik (2013). However, in our algorithm the decision on evaluating or not is made by the position of the particle in the grid (isolated particles are not evaluated) while in the GREEN-PSO the decision is probabilistic and the likelihood of conserving a solution is controlled by a parameter.

The following section gives a formal description of the proposed network and presents the transition rules that define the model for dynamic population structures.

3 PARTIALLY CONNECTED STRUCTURES

Let us consider a rectangular grid G of size $q \times s \geq \mu$, where μ is the size of the population of any population-based metaheuristics or model. Each node G_{uv} of the grid is a tuple $\langle \eta_{uv}, \zeta_{uv} \rangle$, where $\eta_{uv} \in \{1, \dots, \mu\} \cup \{\bullet\}$ and $\zeta_{uv} \in (D \times \mathbb{N}) \cup \{\bullet\}$ for

some domain D . The value η_{uv} indicates the index of the individual that occupies the position $\langle u, v \rangle$ in the grid. If $\eta_{uv} = \bullet$ then the corresponding position is empty. However, that same position may still have information, namely a mark (or clue) ζ_{uv} . If $\zeta_{uv} = \bullet$ then the position is empty and unmarked. Please note that when $q \times s = \mu$, the topology is a static 2-dimensional lattice and when $q \times s = \mu$ and $q = s$ the topology is the standard square grid graph.

In the case of a PSO, the marks are placed by particles that occupied that position in the past and they consist of information about those particles, like their fitness ζ_{uv}^f or position in the fitness landscape, as well as a time stamp ζ_{uv}^t that indicates the iteration in which the mark was placed. The marks have a lifespan of K iterations, after which they are deleted.

Initially, $G_{uv} = (\bullet, \bullet)$ for all $\langle u, v \rangle$. Then, the particles are placed randomly on the grid (only one particle per node). Afterwards, all particles are subject to a movement phase (or grid position update), followed by a PSO phase. The process (position update and PSO phase) repeats until a stop criterion is met.

The PSO phase is the standard iteration of a PSO, comprising position and velocity update. The only difference to a static structure is that in this case a particle may find empty nodes in its neighborhood.

In the position update phase, each individual moves to an adjacent empty node. Adjacency is defined by the Moore neighborhood of radius r , so an individual i at $\rho_g(i) = \langle u, v \rangle$ can move to an empty node $\langle u', v' \rangle$ for which $L_\infty(\langle u, v \rangle, \langle u', v' \rangle) \leq r$. If empty positions are unavailable, the individual stays in the same node. Otherwise, it picks a neighboring empty node according to the marks on them. If there are no marks, the destination is chosen randomly amongst the free nodes.

With this framework, there are two possibilities for the position update phase: stimergeric, whereby the individual looks for a mark that is similar to itself; and Brownian, whereby the individual selects an empty neighbor regardless of the marks. For the first option, let $\mathcal{N}\langle u, v \rangle = \{\langle u^{(1)}, v^{(1)} \rangle, \dots, \langle u^{(w)}, v^{(w)} \rangle\}$ be the collection of empty neighboring nodes and let i be the individual to move. Then, the individual attempts to move to a node whose mark is as close as possible to its own corresponding trait (fitness or position in the fitness landscape, for instance) or to an adjacent cell picked at random if there are no marks in the neighborhood. In the alternative Brownian policy, the individual moves to an

adjacent empty position picked at random. In either case, the process is repeated for the whole population.

For this paper, the investigation is restricted to the Brownian structure. The algorithm is referred in the remaining of the paper as PSO-B, followed by the grid size $q \times s$. An extension of the PSO-B is also proposed by introducing a conservation of function evaluations (*cfé*) strategy. If at a given time-step a particle has no neighbors, then the particle is updated but its position is not evaluated. This version of the algorithm is referred to as PSO-B*cfé*. The following section describes the results attained by the PSOs with dynamic structure and Brownian movement, with and without conservation of function evaluations and compares them to the standard topologies.

4 EXPERIMENTS AND RESULTS

This section describes the experiments conducted for evaluating the performance of the proposed structure. The algorithm is first compared to the version without conservation of function evaluations.

Table 1: Benchmarks for the experiments. Dynamic range, initialization range and stop criteria.

| function | mathematical representation | Range of search/ Range of initialization | stop |
|---------------------|--|---|---------|
| Sphere f_1 | $f_1(\vec{x}) = \sum_{i=1}^D x_i^2$ | $(-100, 100)^{30}$ $(50, 100)^{30}$ | 0.01 |
| Rosenbrock f_2 | $f_2(\vec{x}) = \sum_{i=1}^{D-1} (100(x_{i+1} - x_i)^2 + (x_i - 1)^2)$ | $(-100, 100)^{30}$ $(15, 30)^{30}$ | 100 |
| Rastrigin f_3 | $f_3(\vec{x}) = \sum_{i=1}^D (x_i^2 - 10 \cos(2\pi x_i) + 10)$ | $(-10, 10)^{30}$ $(2.56, 5.12)^{30}$ | 100 |
| Griewank f_4 | $f_4(\vec{x}) = 1 + \frac{1}{4000} \sum_{i=1}^D x_i^2 - \prod_{i=1}^D \cos\left(\frac{x_i}{\sqrt{i}}\right)$ | $(-600, 600)^{30}$ $(300, 600)^{30}$ | 0.05 |
| Schaffer f_5 | $f_5(\vec{x}) = 0.5 + \frac{(\sin \sqrt{x^2 + y^2} - 0.5)}{(1.0 + 0.001(x^2 + y^2))^2}$ | $(-100, 100)^2$ $(15, 30)^2$ | 0.00001 |

Table 2: PSO-B and PSO-B*cfé*. Best fitness values averaged over 50 runs.

| | f_1 | f_2 | f_3 | f_4 | f_5 |
|---------------------------|-------------------------------------|-------------------------------------|-------------------------------------|-------------------------------------|-------------------------------------|
| PSO-B 8×8 | 2.71e-40 ±4.21e-40 | 7.98e+00 ±1.11e+01 | 6.57e+01 ±1.78e+01 | 7.43e-03 ±9.02e-03 | 1.17e-03 ±3.19E-03 |
| PSO-B <i>cfé</i> 8×8 | 3.20e-40 ±8.44e-40 | 1.12e+01 ±2.02e+01 | 6.29e+01 ±1.35e+01 | 7.44e-03 ±7.96e-03 | 1.94e-04 ±1.37E-03 |
| PSO-B 10×10 | 8.74e-38 ±1.29e-37 | 1.25e+01 ±2.07e+01 | 6.23e+01 ±1.88e+01 | 7.73e-03 ±9.50e-03 | 0.00e+00 ±0.00E+00 |
| PSO-B <i>cfé</i> 10×10 | 2.43e-39 ±4.38e-39 | 1.09e+01 ±1.67e+01 | 6.16e+01 ±1.60e+01 | 5.61e-03 ±7.47e-03 | 0.00e+00 ±0.00E+00 |
| PSO-B 15×15 | 3.47e-33 ±4.31e-33 | 1.53e+01 ±2.65e+01 | 6.05e+01 ±1.45e+01 | 3.79e-03 ±6.22e-03 | 3.89e-04 ±1.92E-03 |
| PSO-B <i>cfé</i> 15×15 | 2.47e-45 ±4.21e-45 | 1.31 e+01 ±2.42e+01 | 6.62e+01 ±2.05e+01 | 6.45e-03 ±9.90e-03 | 0.00e+00 ±0.00E+00 |

Then, the topology with Brownian motion and conservation of evaluations is compared to the *lbest*, *gbest*, and standard square *von Neumann* topologies.

An experimental setup was constructed with five benchmark unimodal and multimodal functions that are commonly used for investigating the performance of PSO – see (Kennedy and Mendes, 2002) and (Trelea, 2003) for instance). The functions are described in Table 1. The optimum (minimum) of all functions is located in the origin with fitness 0. The dimension of the search space is set to $D = 30$ (except Schaffer, with $D = 2$). In order to set a square grid graph for the standard *von Neumann* topology, the population size n is set to 49 (which is within the typical range of PSO’s swarm size). The acceleration coefficients were set to 1.494 and the inertia weight is 0.729, as in (Trelea, 2003). $Xmax$ is defined as usual by the domain’s upper limit and $Vmax = Xmax$. A total of 50 runs for each experiment are conducted. *Asymmetrical initialization* is used (the initialization range for each function is given in Table 1).

Two experiments were conducted. Firstly, the algorithms were run for a limited amount of function evaluations (147000 for f_1 and f_5 , 49000 for f_2, f_3 and f_4) and the fitness of the best solution found was averaged over the 50 runs. In the second experiment the algorithms were run for 980000 iterations (corresponding to 20000 iterations of standard PSO with $n = 49$) or until reaching the stop criterion. The criteria were taken from (Kennedy and Mendes, 2002) and are given in Table 1. For each function and each algorithm, the number of function evaluations required to meet the criterion is recorded and averaged over the 50 runs. A success measure is defined as the number of runs in which an algorithm

attains the fitness value established as the stop criterion.

Table 3: PSO-B and PSO-Bcfe. Function evaluations averaged over 50 runs.

| | f_1 | f_2 | f_3 | f_4 | f_5 |
|--------------------------|---|---|--|--|---|
| PSO-B 8×8 | 21070.98 ±1023.60 (50) | 56472.50 ±49342.60 (50) | 13224.90 ±3894.30 (48) | 19959.66 ±1408.36 (50) | 14369.25 ±20599.03 (46) |
| PSO-Bcfe 8×8 | 21157.7 ±1092.81 (50) | 96148.42 ±112675.6 (50) | 13427.12 ±2259.73 (50) | 19968.40 ±1103.20 (50) | 10275.69 ±6952.548 (49) |
| PSO-B 10×10 | 22700.72 ±906.39 (50) | 65769.76 ±70232.15 (50) | 15114.46 ±3939.74 (48) | 21574.70 ±1107.56 (50) | 10741.78 ±10658.16 (50) |
| PSO-Bcfe 10×10 | 21796.04 ±832.54 (50) | 57704.16 ±71260.77 (50) | 13953.04 ±3341.41 (49) | 20430.34 ±1176.32 (50) | 11817.06 ±15647.63 (50) |
| PSO-B 15×15 | 26122.88 ±950.08 (50) | 74321.24 ±83535.59 (50) | 20408.50 ±3692.10 (50) | 24626.42 ±1406.53 (50) | 11830.83 ±11576.37 (48) |
| PSO-Bcfe 15×15 | 19600.76 ±730.62 (50) | 77348.06 ±91374.00 (50) | 16713.59 ±4387.78 (46) | 18734.94 ±1029.26 (50) | 10890.55 ±11624.59 (47) |

Table 4: PSO-Bcfe and standard topologies. Best fitness values averaged over 50 runs.

| | f_1 | f_2 | f_3 | f_4 | f_5 |
|----------------------------|-------------------------------------|-------------------------------------|-------------------------------------|-------------------------------------|-------------------------------------|
| PSO VN | 4.26e-36 ±1.32e-35 | 1.39e+01 ±2.39e+01 | 6.40e+01 ±1.59e+01 | 5.61e-03 ±8.78e-03 | 0.00e+00 ±0.00e+00 |
| PSO <i>lbest</i> | 1.20e-25 ±1.30e-25 | 1.19e+01 ±2.48e+01 | 1.11e+02 ±1.74e+01 | 2.95e-04 ±2.09e-03 | 1.94e-04 ±1.37e-03 |
| PSO <i>gbest</i> | 3.80e+03 ±5.67e+03 | 1.19e+01 ±1.50e+01 | 9.86e+01 ±2.84e+01 | 3.80e+01 ±5.80e+01 | 1.36e-03 ±3.41e-03 |
| PSO-Bcfe 10×10 | 2.43e-39 ±4.38e-39 | 1.09e+01 ±1.67e+01 | 6.16e+01 ±1.60e+01 | 5.61e-03 ±7.47e-03 | 0.00e+00 ±0.00e+00 |

Table 2 shows the average fitness values attained by PSO-B and PSO-Bcfe with different grid sizes. Table 3 shows the average number of function evaluations required to meet the stop criteria as well as the number of successful runs. The performance according to the fitness values is very similar with no significant differences between the algorithm in every function except f_1 (PSO-Bcfe is better). When considering the number of function evaluations (i.e., convergence speed), PSO-Bcfe is significantly better or statistically equivalent in every function. For the statistical tests comparing two algorithms, non-parametric Kolmogorov-Smirnov tests (with 0.05 level of significance) have been used.

The results confirm that PSO-Bcfe is able to improve the convergence speed of PSO-B without

degrading the accuracy of the solutions. The loss of information that results from conserving evaluations is clearly overcome by the benefits of reducing the computational cost per iteration.

In the case of f_1 , PSO-Bcfe also significantly improves the quality of the solutions, namely with larger grids. The proposed scheme seems to be particularly efficient in unimodal landscapes, but further tests are required in order to confirm this hypothesis and understand what mechanisms make PSO-Bcfe so efficient in finding more precise solutions for the sphere function.

Table 5: PSO-Bcfe and standard topologies. Function evaluations averaged over 50 runs.

| | f_1 | f_2 | f_3 | f_4 | f_5 |
|----------------------------|--|---|---|--|---|
| PSO VN | 23530.78 ±954.74 (50) | 72707.18 ±92916.33 (50) | 18424.00 ±11082.75 (49) | 22015.70 ±1304.60 (50) | 17622.36 ±16056.6 8 (50) |
| PSO <i>lbest</i> | 32488.96 ±921.45 (50) | 80547.18 ±112067.67 (50) | 233260.18 ±281453.62 (17) | 30200.66 ±1703.88 1 (50) | 26263.00 ±27266.8 6 (49) |
| PSO <i>gbest</i> | 16082.39 ±2697.41 (33) | 56681.24 ±88165.30 (50) | 9602.04 ±3599.04 (25) | 14856.07 ±2028.12 (27) | 13933.09 ±21576.6 3 (43) |
| PSO-Bcfe 15×15 | 21796.04 ±832.54 (50) | 57704.16 ±71260.77 (50) | 13953.04 ±3341.41 (49) | 20430.34 ±1176.32 (50) | 11817.06 ±15647.63 (50) |

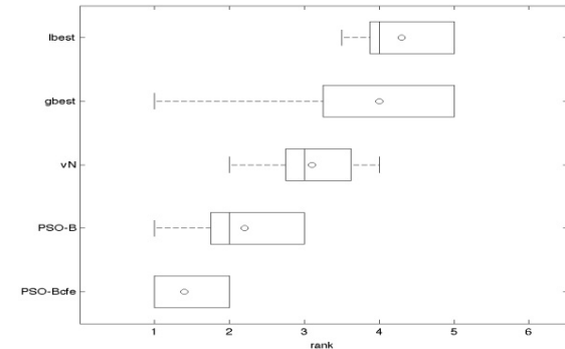


Figure 1: Rank by Holm-Bonferroni test.

Tables 4 and 5 compare the PSO-Bcfe with standard PSOs: square grid (7×7) with von Neumann (VN) neighborhood, *lbest* with $k = 3$ (ring) and *gbest*.

As expected, *gbest* is fast but its solutions are clearly worst than the other algorithms results (except for f_2 , and f_3) and the success rates are lower than the average; *lbest* is fairly accurate when compared to other strategies but it clearly requires more evaluations than the other PSOs in order to meet the

same criteria. The standard *von Neumann* strategy and the PSO-Bcfe are more consistent throughout the test set, but PSO-Bcfe is faster in most of the functions and significantly faster in $f_1, f_3,$ and f_4 . In addition, its solutions for f_1 are significantly better than the fitness values attained by the PSO with *von Neumann* topology. The proposed strategy maintains or improves the accuracy of the *von Neumann* strategy throughout the test set — the consistency of the *von Neumann* topology has been reported by Kennedy and Mendes (2002) —, while increasing significantly the convergence speed.

A statistical analysis of the algorithms (including also PSO-B) on the entire test set was conducted. First, a Friedman test showed that there are no significant differences in the performance of the algorithms considering the fitness values criteria. When taking into account the function evaluations and success rates the test reveals significant differences between the algorithms.

After the Friedman test, a Holm-Bonferroni test was conducted for ranking the algorithms according to their convergence speed and reliability and detect significant differences between the algorithms. PSO-Bcfe ranks first, followed by PSO-B, PSO with *von Neumann* topology, PSO with *lbest* and finally PSO with *gbest* (see Figure 1). Considering $\alpha = 0.1$, PSO-Bcfe is significantly better than the *von Neumann*, *lbest* and *gbest* topologies. Table 6 shows the results of the Holm test.

Table 6: Holm test results. $\alpha = 1$.

| | z-statistic | p-value | α/i |
|------------------|-------------|----------|------------|
| PSO-B | 0.800000 | 0.211855 | 0.10000 |
| PSO VN | 1.700000 | 0.044565 | 0.05000 |
| PSO gbest | 2.600000 | 0.004661 | 0.33333 |
| PSO lbest | 2.900000 | 0.001866 | 0.02500 |

The Holm test concludes that although PSO-Bcfe attains a better average ranking than PSO-B, there are no significant differences between the algorithms. Please remember that the grid size considered in the general comparisons is 10×10 . If we consider a 15×15 grid size we see that PSO-Bcfe is faster in every function and significantly faster in f_1, f_3 and f_4 , while for grids with size 8×8 the algorithms are statistically equivalent in every function. In general, the performance of PSO-Bcfe is improved or preserved when the grid size grows, while the performance of PSO-B degrades with the grid size.

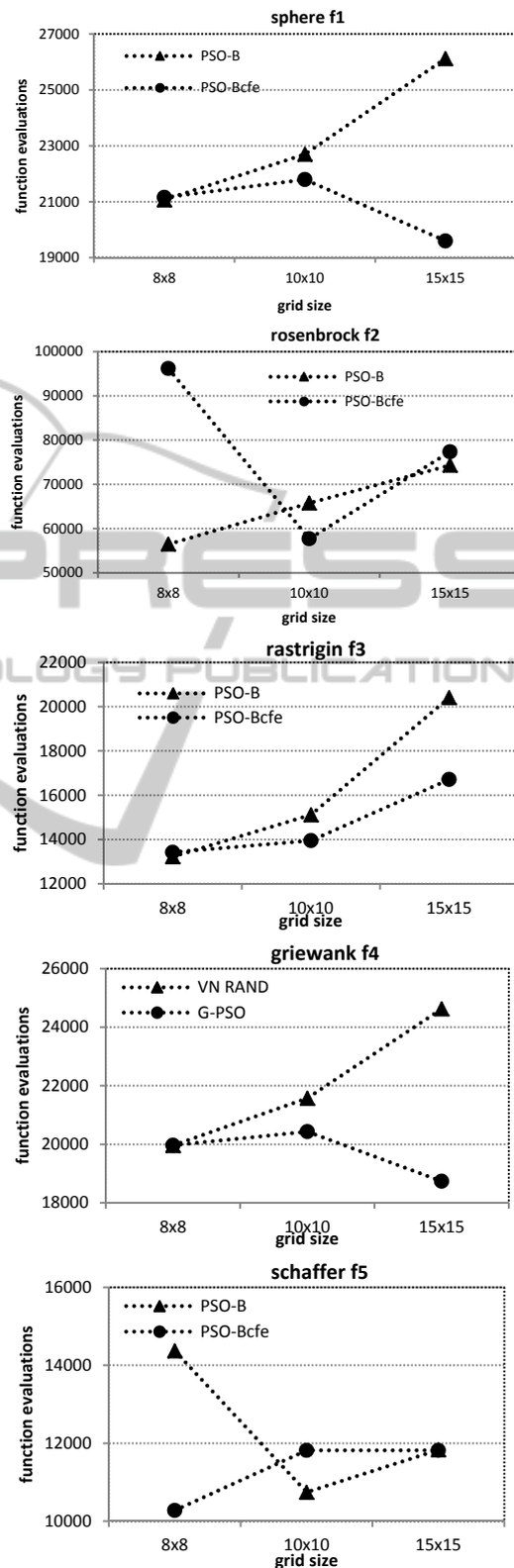


Figure 2: PSO-B and PSO-Bcfe. Function evaluations required to meet stop criteria when using grids with different sizes.

This behavior may be explained by the fact that in larger grids the particles are isolated more often and for longer periods of time. During these periods, the particles are not using information from the rest of the swarm. PSO-Bcfe partially overcomes the loss of communication and information by not evaluating the particles, thus saving computational resources. Figure 2 graphically displays the evaluations required by each version of the algorithm in each function for different grids. Except for f_5 , PSO-B number of function evaluations to meet the criteria increase with the size of the grid, while PSO-Bcfe scales better, namely in functions f_1 , f_3 and f_4 .

These results show that it is possible to improve standard PSO's performance by structuring the particles on a grid of nodes, let them move according to simple rules and save computational resources by letting them follow their current trajectories without evaluation the new positions.

5 CONCLUSIONS

This paper proposes a general scheme for structuring dynamic populations for the Particle Swarm Optimization (PSO) algorithm. The particles are placed on a grid of nodes where the number of nodes is larger than the swarm size. The particles move on the grid according to simple rules and the network of information is defined by the particle's position on the grid and its neighborhood (*von Neumann* vicinity is considered here). If isolated (i.e., no neighbors except itself), a particle is updated but its position is not evaluated. This strategy results in loss of information but it decreases the number of evaluations per generation. The results show that the payoff in convergence speed overcomes the loss of information: the number of function evaluations is reduced in the entire test set, while the accuracy of the algorithm (i.e., the averaged final fitness) is not degraded by the conservation of evaluations strategy.

The proposed algorithm is tested with a Brownian motion rule and compared to standard static topologies. Statistical tests and ranking according to convergence speed and success rates shows that the dynamic structure with conservation of function evaluations ranks first and it is significantly better than the *von Neumann*, ring and *lbest* topologies. Furthermore, the conservation of evaluations strategy results in a more stable performance when varying the grid size, while removing this strategy from the proposed dynamic structure results in a

drop off of the convergence speed when the size of the grid increases in relation to the swarm size.

The present study is restricted to dynamic structures based on particles with Brownian motion. However, a self-organized behavior based on communication via the grid (stigmergy) can be modeled by the general framework proposed in this paper. Future research will be focused on dynamic structures with stigmergic behavior based on the fitness and position of the particles.

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