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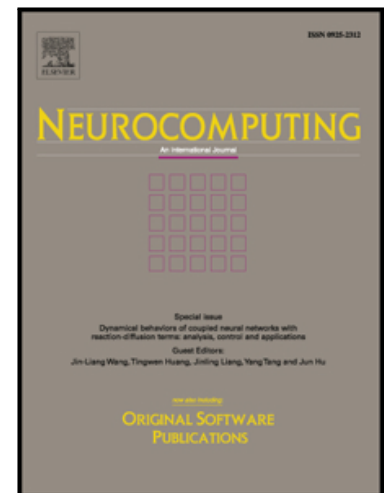
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Parallel population-based algorithm portfolios: An empirical study

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Abstract

Although many algorithms have been proposed, no single algorithm is better than others on all types of problems. Therefore, the search characteristics of different algorithms that show complementary behavior can be combined through portfolio structures to improve the performance on a wider set of problems. In this work, a portfolio of the Artificial Bee Colony, Differential Evolution and Particle Swarm Optimization algorithms was constructed and the first parallel implementation of the population-based algorithm portfolio was carried out by means of a Message Passing Interface environment. The parallel implementation of an algorithm or a portfolio can be performed by different models such as master-slave, coarse-grained or a hybrid of both, as used in this study. Hence, the efficiency and running time of various parallel implementations with different parameter values and combinations were investigated on benchmark problems. The performance of the parallel portfolio was compared to those of the single constituent algorithms. The results showed that the proposed models reduced the running time and the portfolio delivered a robust performance compared to each constituent algorithm. It is observed that the speedup gained over the sequential counterpart changed significantly depending on the structure of the portfolio. The portfolio is also applied to a training of neural networks which has been used for time series prediction. Result demonstrate that, portfolio is

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able to produce good prediction accuracy.

Keywords: Algorithm portfolios, global optimization, neural networks, parallel computing.

1. Introduction

In the last few decades, different kinds of EAs have been developed. These algorithms have shown excellent search abilities but usually their performance may vary considerably according to the problem. No single algorithm performs better than others. Therefore, selection of the most appropriate algorithm for a particular problem needs considerable time. In this case, to solve problems efficiently, researchers use a combination of different algorithms, which is usually referred to as Portfolio Algorithm (PA) [1, 2, 3, 4]. The main aim of these portfolio algorithms is to improve the performance and to make a more stable algorithm by combining the advantages of the optimization algorithms. Although these portfolios provide better results than the single algorithms of the constituting portfolios, their main problem is that their running times to solve large scale problems are unacceptable. Parallel computing systems, which emerged as a result of recent hardware developments, can be employed to overcome this problem. Parallel computing divides tasks into smaller parts and operates the parts on multiprocessor hardware architectures simultaneously. In this way, increased computing needs are met and the solutions are obtained faster. On the other hand, parallel programming has grown even more in popularity because of the availability of multi-core CPUs and it is known that EAs can explore efficiently the use of parallel concepts to speed the process. The main purpose of parallel EAs is to solve a given problem in less time.

Although the portfolio idea is not new, there has not been much research done in parallel implementation of portfolios. A parallel portfolio of algorithms is a collection of different algorithms running on different CPUs. The main aim of the parallel portfolio is not only to enhance the performance of the component algorithms but also to reduce the computation time.

In this study, a Parallel Portfolio Algorithm (PPA) was proposed on different parallelization models through a Message Passing Interface (MPI) [5] among multiprocessing units. We study the possibility of combining algorithms
30 into portfolios and implementing a parallel portfolio including the Artificial Bee Colony (ABC) [6], Differential Evolution (DE) [7] and Particle Swarm Optimization (PSO) [8], which are recent evolutionary computation techniques and of which the speedup performance are analyzed with these parallelization models. Many different versions of these algorithms based on different models have
35 been proposed and used to solve different optimization problems in the literature. Some survey papers have summarized these studies [9, 10, 11, 12] and some papers carried out a review on main metaheuristics and presented their similarities and differences [13].

As mentioned before, no one algorithm outperforms all other algorithms on
40 all types of problems. Some problems that are unsolvable by superior algorithm can be solved with an inferior one. This fact has encouraged researchers to work on portfolio algorithm structures. The earliest portfolio algorithm approach was proposed by Gomes and Selman [1] and Huberman et al. [2] in 1997. These studies applied a portfolio of two or more Las Vegas algorithms to
45 certain combinatorial problems and obtained significant performance improvement. Fukunaga proposed combining different genetic algorithm instances with different control parameter sets and applied them to solve the traveling salesmen problem [3]. Subsequent work focused on showing the computational advantage of a portfolio approach on hard combinatorial search problems [4]. In the following years, a portfolio algorithm for numerical optimization was proposed by
50 Peng et al. [14]. This study emphasized that, selecting the best algorithm is very difficult in a limited budget time and that distributing the time budget to multiple algorithms can reduce this risk. Vrugt and Robinson proposed an adaptive method approach that combine genetic algorithm, evolution strategy and
55 particle swarm optimization in one framework [15]. Some studies that combine a variety of genetic algorithms use population partitioning techniques to obtain better performance [16, 17]. Some studies focused on integrating multiple search

operators or algorithms in a portfolio [18, 19, 20]. Samulowitz et al. developed an algorithm portfolio tool called snappy that is provide a strong baseline and easily extended by its users [21]. More recently, Tang et al. [22] proposed algorithm portfolios based on the Estimated Performance Matrix (EPM-PAP), which is equipped with a novel constituent algorithms selection module. In this paper the proposed model can successfully identify the appropriate constituent algorithm and achieve better results than all the single algorithms considered in their study. Not only selection of the most appropriate algorithm but also their associated control parameter settings for a particular problem needs considerable time. Extensive studies have been done on appropriate setting of the control parameters of EAs [23, 24, 25, 26, 27]. Also some studies focused on adaptive strategies for control parameter selection [28, 29]. Also portfolio algorithm is related to ensemble methods that use multiple learning algorithms to obtain better predictive performance [30, 31]. In ensemble methods while some studies combine different algorithm like algorithm portfolio [32, 33], the others combine same algorithms with different parameters or adaptive strategies [34, 35, 36]. Zhao et al. proposed an ensemble of different neighborhood sizes with online self-adaptation to enhance the multiobjective evolutionary algorithm based on decomposition [37]. Shang et al. proposed a multi-population based cooperative coevolutionary algorithm to solve the multi-objective capacitated arc routing problem [38]. Portfolio algorithm is both effective and robust, but their parallel implementation has received relatively little attention. Petrik and Zilberstein [39] proposed a method for finding a static parallel portfolio of algorithms, in which the share of processor time allocated to each algorithm is fixed. Yun and Epstein [40] proposed an approach that constructs algorithm portfolios for parallel runs based on a combination of a greedy algorithm, and three heuristics. Also portfolio algorithm has been applied to solve various optimization problems from various fields, such as satisfiability problem [41], classification and prediction [42, 43, 44], and scheduling problem [45].

As a result, portfolio algorithms and ensemble methods naturally attract increasing attention. The application of the techniques in previous literature

is aimed to improve the performance of EAs while our study is aimed to not
 90 only on improving the performance of the constituent algorithms but also on
 investigating the PPAs running time in a parallel implementation context. For
 this purpose, we performed different parallelization models such as master-slave
 and coarse-grained and made a comparison among the serial and parallel PPAs.
 The other purpose of the study is to employ the proposed model to a real-
 95 world problem. The real-world problem, training an artificial neural network
 for predicting Mackey Glass and Box Jenkins time series. Although many
 algorithms have been applied successfully for prediction problems [46, 47, 48, 49],
 parallel portfolio algorithms is not available in the literature yet.

The rest of the paper is organized as follows. In Section II, the EAs which
 100 are used in the proposed model are described briefly. Section III provides a
 description of the parallel implementation concept and the proposed parallel
 implementation portfolio is explained. The experiments and results are pre-
 sented in Section IV. Section V contains some conclusions.

2. Algorithms

105 Some recent EAs include ABC, DE and PSO. A brief description of these
 algorithms is presented in the following paragraphs.

2.1. Artificial bee colony

The ABC algorithm was introduced by Karaboga in 2005 and is inspired
 by the foraging behavior of honey bees [6]. The working process of the ABC
 110 algorithm contains three different phases. The first phase is that of the employed
 bees which are responsible for exploiting all the sources. The second phase is
 that of the onlooker bees which use potentially rich sources. The third phase is
 that of the scout bees which are responsible for exploring undiscovered sources.
 “Limit”, which is a parameter of the ABC algorithm, uses exhausted sources. In
 115 the ABC algorithm, employed bees and onlooker bees use the same perturbation
 strategy which is defined by Eq. (1).

$$x_{ij}(t+1) = x_{ij}(t) + rand()(x_{ij}(t) - x_{r_1j}(t)) \quad (1)$$

where r_1 is the randomly selected solution vector ($r_1 \neq i$) and j is the randomly selected index. The $rand()$ function generates a uniformly distributed random variable $\in [0, 1]$. Detailed information on the ABC algorithm can be
 120 found in [6, 50].

2.2. Differential evolution

The DE algorithm was introduced by Storn and Price in 2005 [7]. It uses different recombination operators to improve the population. In this paper we use the DE/rand/1/bin configuration. This configuration's update formulas are
 125 given by Eqs. (2) and (3).

$$v_i(t) = x_{r_1}(t) + F(x_{r_2}(t) - x_{r_3}(t)) \quad (2)$$

$$x_i(t+1) = \begin{cases} v_i(t) & \text{if } rand() \leq CR \\ x_i(t) & \text{else} \end{cases} \quad (3)$$

The weighting factor $F \in [0, 2]$ controls the amplification of differential variation. The crossover rate $CR \in [0, 1]$ probabilistically controls the amount of recombination. x_{r_1} , x_{r_2} and x_{r_3} are randomly selected solution vectors ($r_1 \neq r_2 \neq r_3 \neq i$). The $rand()$ function generates a uniformly distributed random
 130 variable $\in [0, 1]$. Detailed information on the DE algorithm can be found in [7].

2.3. Particle swarm optimization

The PSO algorithm, which was introduced by Kennedy et al. in 1995, inspired by social and cooperative behavior displayed by various species like birds, fish etc [8]. It modifies velocities based on personal best and global best. The
 135 modification of velocities and positions are obtained by using Eqs. (4) and (5).

$$x_{id} = x_{id} + v_{id} \quad (4)$$

$$v_{id} = wv_{id} + c_1r_1(x_i^{pb} - x_{id}) + c_2r_2(x^{gb} - x_{id}) \quad (5)$$

where x_{id} is the i^{th} particle's position and v_{id} is the velocity for the i^{th} particle. x^{pb} and x^{gb} are the personal best and global best positions respectively. w is the inertia weight, which determines how the previous velocity of the particle influences the velocity in the next iteration. c_1 and c_2 are acceleration constants. r_1 and r_2 are the uniformly generated random number in the range of $\in [0, 1]$. Detailed information on the PSO algorithm can be found in [8].

3. Parallel models of portfolio algorithms

Parallel EAs can be performed based on different models which are categorized depending on the synchronization and the communication behaviors of subtasks. In this section, we briefly introduce parallel EA models and discuss the differences between them. In general, there are three different well-known models for the parallelization of EAs which was named master-slave, fine-grained and coarse-grained [51]. These models are shown in Figure 1.

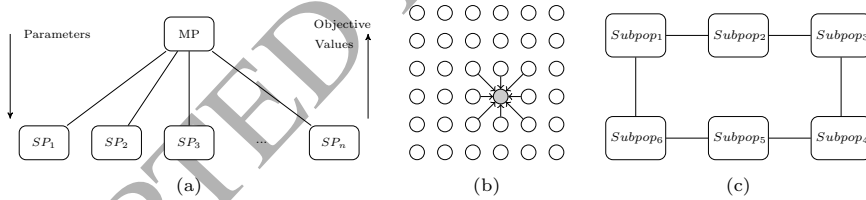


Figure 1: Parallel computing models, a)Master-slave, b)Fine-grained, and c)Coarse-Grained (MP:Master processor, SP:Slave Processor)

In the master-slave parallelization approach, while the evaluation process of each individual in the population is performed on a different slave processor, the selection and recombination processes take place in the master processor as given in Figure 1(a). The fitness value of each individual is independent of the others and there is no communication during the evaluation of the objective function. After evaluation, the slave processor sends its results to the master processor and the master processor performs the selection and recombination operations

to create offsprings. The offsprings are distributed to slave processors to be assigned a fitness value, and the process continues in this way. Communication occurs only when each slave sends the fitness value and receives its subset of individuals to evaluate.

160 In the fine-grained parallelization model, the population is distributed over the processors. Parallelism and communication are realized among some individuals a local neighborhood structure as given in Figure 1(b). The improvement when more processors used is limited. Using more processors for a small problem increases the running time significantly as a result of excessive communication
165 between processors. That is, an individual can only compete and mate with its nearby neighbors. The topology of the neighborhood and the number of individuals in the neighborhood affects the performance of the parallel algorithm. This model is initially designed for working in massively parallel machines.

In the coarse-grained parallelization model the algorithms are executed con-
170 currently on several independent subpopulations and they regularly exchange some individuals with those from other subpopulations during their search as given in Figure 1(c). In this model, new control parameters such as the number of individuals to migrate, the migration interval that controls the migration frequency and the topology which defines the neighbors of each island need to
175 be set carefully.

Many hybrids have been defined by combining the above models, such as coarse-grained and fine-grained, coarse-grained and master-slave, and coarse-grained at two levels. The hybrid approach allows the advantages of two models to be combined.

180 All parallelization models have advantages and disadvantages. Parallel implementation of EAs may be complex and there are some problems with regard to selecting the parallel computing model and its optimum parameter setting.

In the master-slave model, the algorithms' search behavior does not change, it is exactly like a serial. As with other models, this model does not require addi-
185 tional parameters. This model is most suitable whenever the fitness evaluations are significantly expensive. The communication overhead between the master

processor and the slave processors becomes significant. The fine-grained model is designed for working in massively parallel machines and it can easily parallelize many features of EAs such as selection, mating, survival etc. The speedup and performance of the algorithms may significantly change the models' parameters: topology, neighborhood size and complicated shape. The coarse-grained model is easy to implement. When the fitness function is computationally inexpensive, this model is probably the most preferred. It needs less communication between nodes, so its efficiency may be better than those of other models. However, when EAs are parallelized based on the coarse-grained model, more control parameters are introduced to the approach, as mentioned before. The speedup and performance of the algorithms based on the coarse-grained model may significantly change due to the differences in the algorithms' behaviors. Therefore, additional control parameters need to be set carefully. Hybrid models generally to achieve the combined advantages of two models and give better performance than any of them alone.

3.1. Parallel population-based algorithm portfolios

A parallel portfolio of algorithms is a collection of different algorithms running on parallel computing systems. Their parallel implementation can be performed by different models. In this study, we explored two types of these models: coarse-grained and a hybrid model.

In our first model, the coarse-grained parallelization model was used to implement the parallel portfolio algorithm, in which each node runs the different algorithms independently and is initialized by its own set. Migration of individuals among sub-algorithms is performed in a certain interval to encourage the exchange experience of each independent algorithm among the sub-algorithms. The migration scheme adopted in the study uses a global sorting step in which the worst individual in m th sub-algorithms is replaced with the best individual of each of the $(m - 1)$ th sub-algorithms. The pseudo code of the PPA based on the coarse-grained parallelization model is shown in Algorithm 1.

In our second model, the hybrid parallelization model was used to imple-

Algorithm 1 Pseudo code of the PPA based on the coarse-grained parallelization model

```

1: Initialize algorithms' parameters
2: Initialize each of sub-algorithms of solutions;
3: Evaluate each population in  $pop_{ABC}$ ,  $pop_{DE}$ ,  $pop_{PSO}$ ;
4: Counter =1;
5: while  $Counter \neq MaximumCycleNumber$  do
6:   for each of sub-algorithms do                                     ▷ Begin parallel block
7:     if  $Counter(mod\ frequency) = 0$  then
8:       Send the best ind. to the next neighbour (MPI_Send);
9:       Receive the best ind. from the previous neighbour (MPI_Recv);
10:      Replace individuals in the population;
11:    end if
12:    Run each sub-algorithm with new population;
13:    Evaluate solutions in the populations;
14:    Memorize the best solution;
15:  end for                                                         ▷ End parallel block
16:  Counter = Counter +1
17: end while

```

ment the two level parallel portfolio algorithm. In this hybrid model, the upper level it is coarse-grained, and the lower level it is master-slave. At the upper level each node runs the different algorithms independently, and at the lower level the master processor sends the offspring to the slave processors and performs selection/updates only. The pseudo code of the PPA based on the hybrid parallelization model is shown in Algorithm 2.

Algorithm 2 Pseudo code of the PPA based on the hybrid parallelization model

```

1: Initialize algorithms' parameters
2: Initialize each of sub-algorithms of solutions;
3: Evaluate the objective function in parallel (Alg. 3.);
4: Counter =1;
5: while  $Counter \neq MaximumCycleNumber$  do
6:   for each of sub-algorithms do ▷ Begin parallel block
7:     if  $Counter(mod\ frequency) = 0$  then
8:       Send the best ind. to the next sub-algorithm (MPI.Send);
9:       Receive the best ind. from the previous sub-algorithm (MPI.Recv);
10:      Replace individuals in the population;
11:    end if
12:    Run each sub-algorithm with new population;
13:    Evaluate the objective function in parallel (Alg. 3.);
14:    Memorize the best solution;
15:  end for ▷ End parallel block
16:  Counter = Counter +1
17: end while

```

4. Simulation Results

4.1. The experimental environment

The computer platform used to perform the experiments was based on AMD Opteron 6172 with the number of 12 CPU cores and 128 GB 1333 DDR3 RAM computers with a scientific Linux operating system at TUBITAK ULAKBIM

Algorithm 3 Evaluation of fitness in parallel

```

1: for all the slaves process do
2:   Send an individual to a slave process (MPI.Send);
3: end for
4: while individual without fitness value do
5:   Receive result from a slave process (MPI.Recv);
6:   Send an individual to a slave process (MPI.Send);
7: end while
8: for all the slaves process do
9:   Receive result from a slave process (MPI.Recv);
10: end for

```

High Performance Computing Center. The proposed algorithm was implemented in C++ using the MPI library.

230 MPI is a communication protocol for the development of wide variety of parallel computers. MPI's goals are high performance and scalability on all platform. So it provides a platform independent message transfer standards. In this study basic MPI functions were used. The functions MPI.Init and MPI.Finalize are used to initiate and shut down an MPI computation, respectively. The functions MPI.Send and MPI.Recv are used to send and receive a message, respectively. At last, the MPI.Comm.Size is used determine the number of processes in a computation and the MPI.Comm.Rank is used determine the identifier of current process.

4.2. The benchmark suite

240 The proposed algorithms were tested on twenty CEC2010 special session benchmark functions [52]. All functions were given for the special case of dimension $D=1000$. The test suite includes separable functions, partially separable functions, partially separable functions that consist of multiple independent subcomponents, each of which is m-nonseparable, and fully nonseparable functions.

4.3. Experimental settings

In the experiments, synchronous versions of the basic ABC, DE and PSO algorithms were used. The values of the same parameters used in each algorithm, such as population size and maximum number of evaluations, were chosen to be the same as specified in competition [52]. Population size was 100 and the maximum number of function evaluations was 3.000.000; all the runs were repeated 30 times to compare the algorithms statistically. In ABC, the limit was set to 1000. In DE, strategy was set to DE/rand/1/bin, the weighting factor was set to 0.5 and crossover weight was set to 0.3 as recommended in [23, 24, 25]. In PSO, the learning factors were both set to 1.8 and inertia weight was set to 0.6 as recommended in [26]. Table 1 presents the distribution of the population sizes for all PPA instantiations.

Table 1: Sizes of subpopulations allocated to the constituent algorithms of each instantiation of PPA

Algorithm	Algorithm	ABC	DE	PSO
PPA_{21}	ABC+DE	50	50	-
PPA_{22}	ABC+PSO	70	-	30
PPA_{23}	ABC+PSO	40	-	60
PPA_{24}	DE+PSO	-	70	30
PPA_{25}	DE+PSO	-	40	60
PPA_{31}	ABC+DE+PSO	26	50	24
PPA_{32}	ABC+DE+PSO	30	30	40

4.4. Performance metrics

Various methods and metrics are used to measure the performance of a certain parallel algorithm. Some performance metrics, such as speedup and efficiency are most frequently used to measure the performance of a parallel algorithm. Speedup is the ratio of sequential execution time to parallel execution time (Eq. 6) [53]. The optimal value for the speedup is equal to the number of processors. Efficiency is the ratio of speedup to the number of processors (Eq. 7) and the ideal value for it is 1 [53].

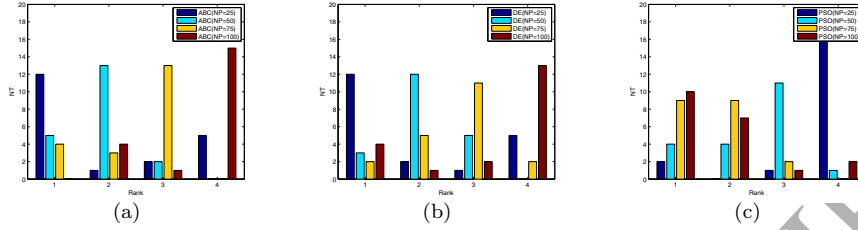


Figure 2: The performance ranking of serial algorithms with a different number of population
a) ABC, b) DE and c) PSO (NT: The number of times the algorithm ranked first)

$$\text{speedup} = \frac{\text{execution time on one processor}}{\text{execution time on } m \text{ processor}} \quad (6)$$

$$\text{efficiency} = \frac{\text{speedup}}{m} \quad (7)$$

4.5. Effect of the population size to the algorithms

Firstly, we analyzed which size of population is better for the constituent algorithms' serial models on CEC2010 test functions. For this purpose, the population size is investigated for four different cases: 25, 50, 75 and 100. The experimental results are given in Table 2 for the ABC, DE and PSO algorithms. Also, to be able to make good a comparison, the performance ranking of serial algorithms is given in Figure 2.

From Table 2 and Figure 2, it can be concluded that as the population size increases, the PSO algorithm generally produces better results while ABC and DE algorithms generally produces worse results. For the test problems employed in this work, the population size of 75–100 can provide acceptable results for PSO and 25–50 can provide acceptable results for ABC and DE algorithms. This experiment helps to decide the optimum individual number of the population in the PPA.

4.6. Coarse-grained PPA

In the second experiment, we analyzed coarse-grained PPA combinations of two and three constituent algorithms. The statistical indicators (mean and

Table 2: Function error values of the solutions obtained by ABC, DE, PSO for the benchmark functions

Function	Population Size											
	25			50			75			100		
	ABC	DE	PSO	ABC	DE	PSO	ABC	DE	PSO	ABC	DE	PSO
f_1	3.64E-14	6.30E+04	1.21E+10	3.34E-14	2.29E+02	9.19E+09	6.96E-09	1.07E+02	9.47E+09	2.52E-06	3.45E+01	9.35E+09
f_2	7.77E-01	3.56E+00	8.63E+03	8.29E-01	2.34E-01	8.12E+03	2.09E+01	6.75E-02	7.84E+03	7.12E+01	1.27E-02	7.67E+03
f_3	2.09E-12	2.39E-01	1.96E+01	4.40E-10	1.03E-01	1.95E+01	1.28E-05	3.89E-02	1.95E+01	1.45E-03	2.21E-02	1.95E+01
f_4	4.17E+13	5.30E+13	4.51E+12	4.01E+13	6.67E+13	3.46E+12	4.10E+13	7.56E+13	2.57E+12	4.04E+13	8.56E+13	3.02E+12
f_5	6.00E+08	5.13E+08	3.60E+08	5.85E+08	5.02E+08	3.41E+08	5.62E+08	5.21E+08	3.27E+08	5.66E+08	5.37E+08	3.16E+08
f_6	1.99E+07	1.97E+07	1.07E+07	1.99E+07	2.01E+07	6.48E+06	2.00E+07	2.03E+07	4.23E+06	2.01E+07	1.97E+07	4.31E+06
f_7	3.81E+10	4.73E+10	3.67E+10	4.24E+10	5.84E+10	2.61E+10	4.34E+10	6.56E+10	1.95E+10	4.65E+10	6.89E+10	1.49E+10
f_8	7.64E+06	2.29E+07	1.63E+14	5.22E+06	2.06E+07	6.73E+13	5.86E+06	3.23E+07	2.89E+13	1.58E+07	5.74E+07	7.31E+13
f_9	3.60E+08	9.37E+08	6.94E+09	4.84E+08	1.26E+09	6.61E+09	6.02E+08	1.52E+09	6.10E+09	7.17E+08	1.74E+09	5.98E+09
f_{10}	7.24E+03	6.50E+03	8.98E+03	7.12E+03	6.89E+03	8.53E+03	7.17E+03	7.13E+03	8.32E+03	7.27E+03	7.41E+03	8.22E+03
f_{11}	2.00E+02	2.04E+02	2.14E+02	2.01E+02	2.05E+02	2.13E+02	2.01E+02	2.05E+02	2.13E+02	2.02E+02	2.07E+02	2.12E+02
f_{12}	3.64E+05	8.90E+05	1.26E+06	5.23E+05	1.08E+06	1.05E+06	6.39E+05	1.22E+06	8.37E+05	7.38E+05	1.32E+06	9.16E+05
f_{13}	6.40E+02	1.56E+04	1.24E+10	4.72E+02	7.69E+03	1.26E+10	4.65E+02	1.85E+03	8.16E+09	6.32E+02	2.62E+03	8.59E+09
f_{14}	8.05E+08	2.71E+09	1.74E+09	1.16E+09	3.69E+09	1.45E+09	1.49E+09	4.48E+09	1.66E+09	1.82E+09	5.04E+09	1.67E+09
f_{15}	1.47E+04	1.40E+04	9.13E+03	1.46E+04	1.48E+04	8.64E+03	1.45E+04	1.53E+04	8.47E+03	1.46E+04	1.58E+04	8.51E+03
f_{16}	3.99E+02	4.10E+02	3.89E+02	4.01E+02	4.14E+02	3.89E+02	4.03E+02	4.17E+02	3.89E+02	4.05E+02	4.16E+02	3.89E+02
f_{17}	6.93E+05	1.94E+06	1.22E+06	1.03E+06	2.41E+06	1.42E+06	1.26E+06	2.70E+06	1.05E+06	1.46E+06	2.92E+06	1.12E+06
f_{18}	1.04E+04	2.12E+04	4.95E+10	7.74E+03	4.50E+03	4.08E+10	6.55E+03	6.46E+03	3.56E+10	7.11E+03	1.00E+04	3.37E+10
f_{19}	5.72E+06	5.03E+06	1.92E+06	6.88E+06	8.60E+06	2.09E+06	7.80E+06	1.06E+07	2.37E+06	8.54E+06	1.22E+07	2.53E+06
f_{20}	1.22E+01	3.31E+04	5.37E+10	1.49E+01	3.38E+03	4.85E+10	8.50E+01	2.78E+03	4.34E+10	5.33E+02	3.48E+03	4.31E+10

standard deviation values) and average running time for 30 runs of the ABC, DE, PSO and all PPA instantiations are given in Table 4. The performance ranking of the coarse-grained PPA and its two and three constituent algorithms are given in Figures 3 and 4 respectively. Running time comparison between the coarse-grained PPA and its constituent algorithms on the 5 test functions with shortest running time (f_3 , f_7 , f_8 , f_{13} , f_{20}) and 5 test functions with longest running time (f_9 , f_{14} , f_{15} , f_{16} , f_{19}) for all PPA instantiations are given in Figures 5- 8.

The portfolio of the basic models of the ABC, DE and PSO algorithms does not achieve the best performance on the CEC2010 functions which take so much time to solve. However, each constituent algorithm delivers the best performance on some problems while it tends not to be successful on others,

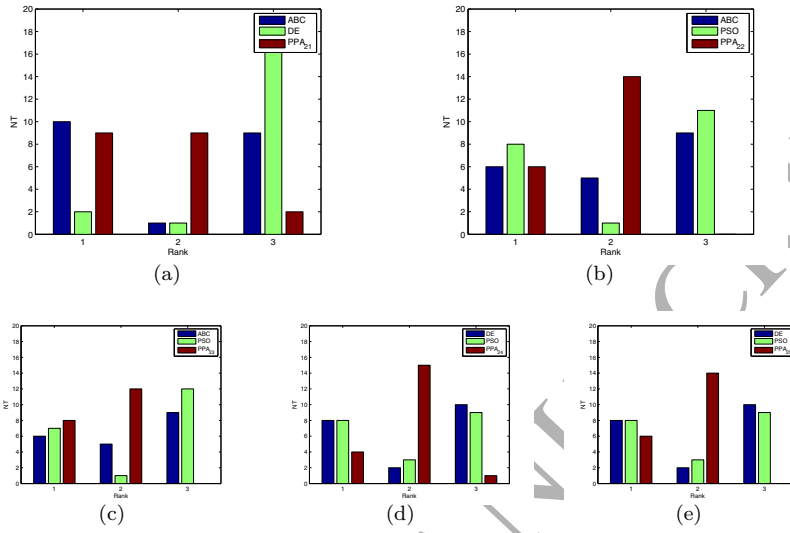


Figure 3: The performance ranking of coarse-grained PPA and its 2 constituent algorithms a) PPA_{21} , b) PPA_{22} , c) PPA_{23} , d) PPA_{24} and e) PPA_{25} (NT: The number of times the algorithm ranked first)

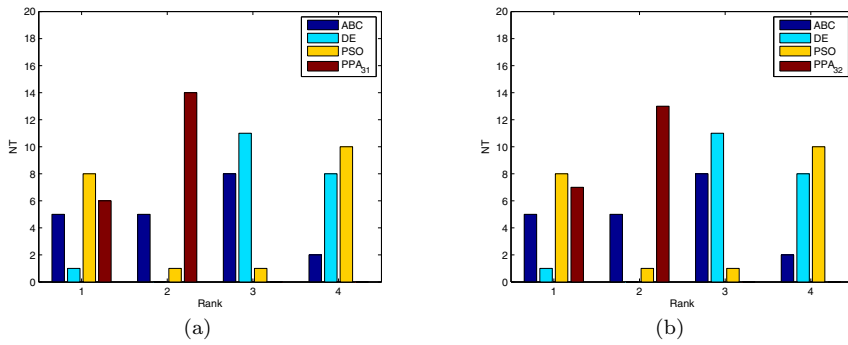


Figure 4: The performance ranking of coarse-grained PPA and its 3 constituent algorithms a) PPA_{31} and b) PPA_{32} (NT: The number of times the algorithm ranked first)

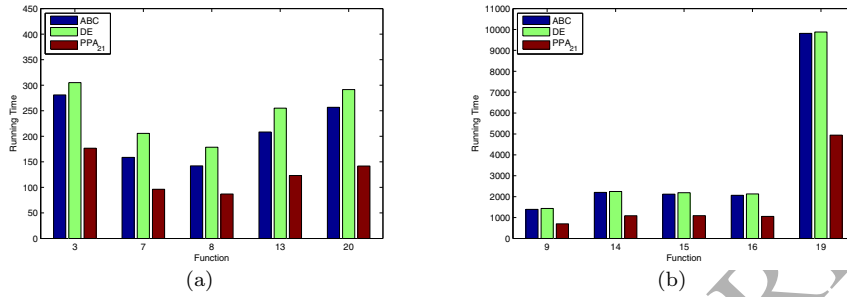


Figure 5: Running time (s) comparison between the coarse-grained PPA_{21} and its 2 constituent algorithms on the 10 test functions a) 5 test functions with shortest running time (f_3 , f_7 , f_8 , f_{13} , f_{20}), b) 5 test functions with longest running time (f_9 , f_{14} , f_{15} , f_{16} , f_{19})

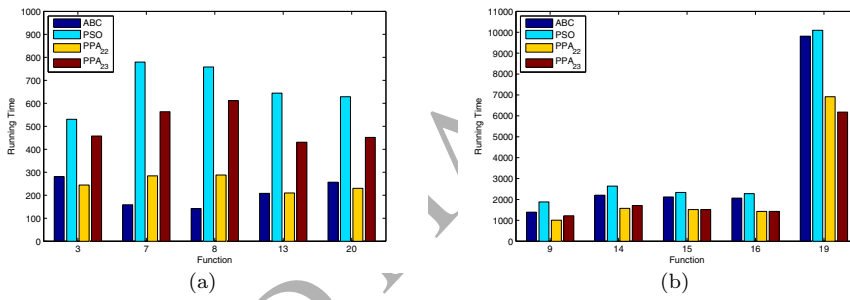


Figure 6: Running time (s) comparison between the coarse-grained PPA_{22} , PPA_{23} and its 2 constituent algorithms on the 10 test functions a) 5 test functions with shortest running time (f_3 , f_7 , f_8 , f_{13} , f_{20}), b) 5 test functions with longest running time (f_9 , f_{14} , f_{15} , f_{16} , f_{19})

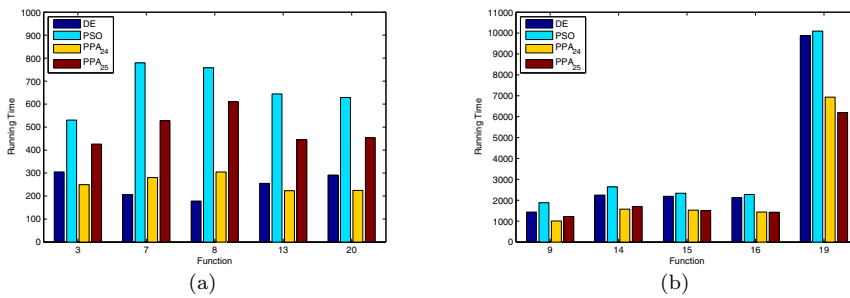


Figure 7: Running time (s) comparison between the coarse-grained PPA_{24} , PPA_{25} and its 2 constituent algorithms on the 10 test functions a) 5 test functions with shortest running time (f_3 , f_7 , f_8 , f_{13} , f_{20}), b) 5 test functions with longest running time (f_9 , f_{14} , f_{15} , f_{16} , f_{19})

Table 3: Running time values obtained by ABC, DE, PSO for the benchmark functions

Function	Population Size											
	25			50			75			100		
	ABC	DE	PSO	ABC	DE	PSO	ABC	DE	PSO	ABC	DE	PSO
f_1	512.71	534.02	1164.26	482.41	518.13	1022.68	486.13	537.05	1002.39	481.53	526.45	993.62
f_2	319.94	301.09	891.06	325.06	339.81	630.72	347.53	374.00	605.79	355.50	378.64	600.57
f_3	223.12	264.95	850.30	235.73	270.44	570.51	265.76	297.38	537.20	280.96	304.99	530.67
f_4	692.27	713.19	1487.05	662.38	705.00	1284.56	668.50	741.51	1200.27	661.57	718.28	1160.67
f_5	532.14	535.89	1110.68	516.31	546.35	856.97	537.12	581.55	828.27	541.72	579.52	817.82
f_6	445.38	455.53	1062.10	432.77	467.30	898.30	455.52	506.42	865.19	467.33	505.61	848.53
f_7	163.91	202.20	968.50	157.94	201.30	869.31	159.95	217.88	816.28	158.61	205.73	779.34
f_8	145.19	183.74	958.73	139.89	182.47	863.69	142.66	183.50	813.20	142.10	178.56	758.06
f_9	1440.27	1431.61	2023.72	1388.95	1422.52	1930.11	1411.86	1429.26	1908.85	1392.27	1436.02	1882.60
f_{10}	1318.60	1304.78	1808.96	1275.28	1310.63	1556.91	1303.40	1318.72	1529.69	1292.82	1337.41	1521.97
f_{11}	1231.24	1236.26	1719.37	1199.81	1245.93	1468.96	1230.58	1249.32	1453.20	1227.34	1277.74	1452.88
f_{12}	397.47	426.79	909.29	383.65	427.16	793.23	392.22	423.19	762.64	384.40	437.88	767.29
f_{13}	211.37	248.04	835.18	207.00	243.59	686.43	212.17	241.36	660.46	208.32	255.08	644.27
f_{14}	2283.64	2232.15	2819.67	2199.11	2245.18	2692.04	2235.92	2242.65	2662.26	2202.63	2249.56	2641.05
f_{15}	2202.55	2155.61	2610.22	2119.90	2169.39	2371.22	2152.96	2186.23	2343.24	2119.82	2190.69	2335.41
f_{16}	2137.39	2097.79	2432.28	2057.04	2106.34	2273.50	2091.12	2117.13	2269.39	2064.19	2129.64	2276.08
f_{17}	657.10	677.95	1211.36	631.86	679.24	1071.23	642.05	680.81	1038.45	633.04	697.33	1036.42
f_{18}	298.97	317.91	815.54	284.15	323.72	660.30	286.29	328.58	654.12	300.52	326.66	648.41
f_{19}	10219.82	9853.06	10234.31	9808.20	9852.01	10125.19	9962.01	9859.02	10123.75	9811.42	9878.49	10093.88
f_{20}	267.68	284.20	815.48	255.50	285.67	653.41	265.12	288.18	630.50	256.80	291.38	628.80

295 while the portfolio delivers a robust performance. The results in Table 4 and performance ranking figures in Figures 3 and 4 show that, in most of the cases, the portfolio is in rank 1 or 2 while the constituent algorithms ranking changes according to the function considered. These results indicate that all proposed PPA instantiations have a robust performance.

300 It was seen that each algorithm has different running times and the running time of the PSO algorithm in large scale problems is much more compared to DE and ABC due to its perturbation operator which changes all parameters, while those the others change only some parameters. Distribution also has effect on both the performance and the running time of the portfolio. If more individuals
305 are assigned to the fastest algorithm while less individuals are assigned to the slower ones, a gain is obtained in the running time of the portfolio, as expected.

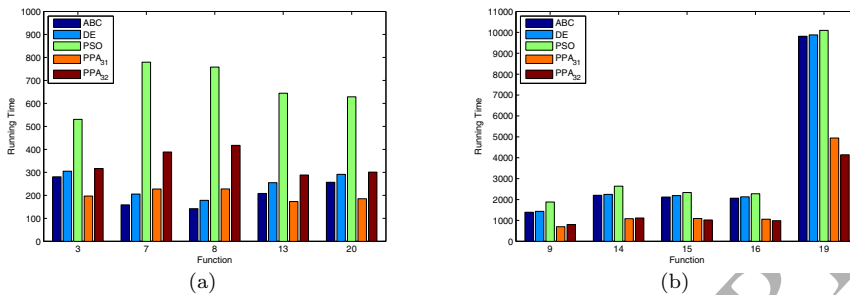


Figure 8: Running time (s) comparison between the coarse-grained PPA_{31} , PPA_{32} and its 3 constituent algorithms on the 10 test functions a) 5 test functions with shortest running time (f_3 , f_7 , f_8 , f_{13} , f_{20}), b) 5 test functions with longest running time (f_9 , f_{14} , f_{15} , f_{16} , f_{19})

However, assigning more individuals to the fastest algorithm may not always yield the best performance in terms of solution quality.

The speedup in the parallel implementation of the coarse-grained model is expected to be linear since a CPU communicates to only one other CPU and then there is not much communication between CPUs as compared to other models (master-slave, fine-grained and hybrid). The linearity of the running time of a portfolio is measured by the running time of the portfolio versus the constituent algorithms. Because the constituent algorithms have different running times, the linearity of the speedup of the subpopulation varies depending on the control algorithm. If the control algorithm is chosen as the slowest algorithm, the speedup is linear while the speedup is less when one of the other algorithms is chosen as the control algorithm.

4.7. Hybrid PPA

In this section, we analyzed the hybrid PPA with respect to the running times. Because of the proposed model includes three different algorithms and each algorithm needs at least two CPUs such that one is master and one is slave, the minimum number of cores for proposed model can be 6. To test the efficiency of the parallelism, the PPA is executed on 6, 9, 12 and 24 processors. The average running times of the hybrid PPA are given in Table 5. Figure 9

show the graph of the parallel efficiency results based on PSO algorithm on the 5 test functions with the shortest running time ($f_3, f_7, f_8, f_{13}, f_{20}$) and the 5 test functions with the longest running time ($f_9, f_{14}, f_{15}, f_{16}, f_{19}$) for PPA_{31} .

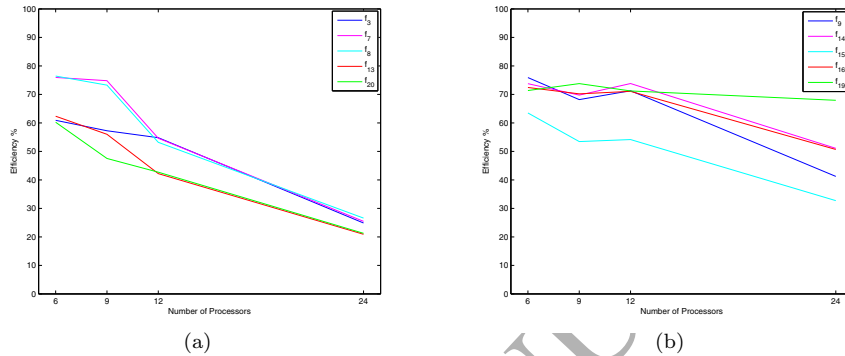


Figure 9: Efficiency result for hybrid PPA_{31} algorithm based on PSO algorithm on the 10 test functions a) 5 test functions with shortest running time ($f_3, f_7, f_8, f_{13}, f_{20}$), b) 5 test functions with longest running time ($f_9, f_{14}, f_{15}, f_{16}, f_{19}$)

Because the hybrid model is a composition of both the coarse-grained and master-slave models, the performance depends on all the factors by which each model is affected, such as the running time of constituent algorithms, distribution of the population and the number of CPUs. The average running times in Table 5 and parallel efficiency graph in Figure 9 show that the efficiency in this model is less for functions with low computation times ($f_3, f_7, f_8, f_{13}, f_{20}$) and much more for functions with high computation times ($f_9, f_{14}, f_{15}, f_{16}, f_{19}$). The average parallelization efficiency is about 70% for the f_{19} function, which has the highest computation cost. Also, increasing the number of CPUs dramatically provides no more gain and even causes the parallelization efficiency to decrease, especially in cases with shortest running times. These results clearly show that the function evaluation time and communication between the processors play an important role. Using more processors for a small problem increases the running time as a result of excessive communication between the processors. Thus we can say that the hybrid PPA algorithm can achieve a

reasonable speedup for expensive fitness functions with a moderate number of
 345 processors.

4.8. PPA for Time Series Prediction Problems

In the last experiment, the performance of the PPA algorithm was applied
 on two well-known time series prediction problems: Mackey Glass (MG) time
 series and Box-Jenkins (BJ) time series which have been in literature to compare
 350 the performance of different neural networks and neurofuzzy systems.

The MG time series is a sequence produced by a nonlinear time delay differ-
 ential equation [54]. The task of this benchmark is to predict the value of the
 time series at the point $(t + 6)$ from the earlier points $(t - 18)$, $(t - 12)$, $(t - 6)$
 and (t) . The BJ time series was recorded from a combustion process of methane
 355 air mixture [55]. The input $u(t)$ is the gas flowing rate and the output $y(t)$ is
 CO_2 concentration. In most of the studies, it has been stated that the best set
 of input variables for predicting $y(t)$ is $y(t - 1)$ and $u(t - 4)$, which is used in
 our study.

In this study, log-sigmoid transfer function, Mean Squared Error (MSE)
 360 performance function and seven network structures that include one hidden
 layer MLP networks with 2,4,6,8,12,16 and 20 hidden nodes were used. The
 maximum generation was set to 1000 and the other parameters of PPA were
 set to default values in experimental settings. The data were normalized to
 the range $[0, 1]$ and shuffled randomly before training of the network. All the
 365 experiments are implemented for 30 runs. The neural networks were performed
 using 450 training data, 500 testing data points for MG time series and 140
 training data, 150 testing data points for BJ time series. In Table 6 average
 MSE values of the neural networks with different structures trained using the
 PPA algorithm are presented. The performance of the PPA were compared
 370 with back propagation (BP), PSO, cooperative random learning particle swarm
 optimization (CRPSO), and genetic algorithm (GA) results data is from recently
 reported publication [56] in Table 7.

Table 6 shows that on the prediction problems, the MPL that showed the

best performance is the topology using 12 hidden nodes for MG and 8 hidden
375 nodes for BJ time series. From Table 7, it is observed that in both training
and testing cases PPA performed better than the other algorithms used in our
study. So, it can be concluded that PPA is effective for time series problems.

5. Conclusions

We have proposed a method for enhancing the robustness of optimization
380 algorithms by running more algorithms in parallel to solve the same optimization
problems. For this purpose, we constructed the PPA with basic models of the
ABC, DE and PSO algorithms. The PPA does not guarantee the best results.
However, each constituent algorithm delivers the best performance on some
problems while it tends not to be successful on others, while the PPA delivers
385 a robust performance. The results clearly showed that the PPA has advantages
when a number of different problems need to be solved. Also, the performance
of the PPA is quite sensitive to the distribution of the constituent algorithms.
However, description of any one portfolio which is the best, is very difficult. In
particular, if you use more constituent algorithms in a PPA, finding the optimal
390 PPA is not possible.

We explored two types of parallel implementation model: coarse-grained
and a hybrid model. These models are very suitable for portfolio structures.
Empirical studies demonstrated that each implementation of the PPA gained
speedup when compared to the sequential counterparts. In addition, we explored
395 how to affect the constituent algorithms' distribution in the population in terms
of running time. The result clearly showed that, distribution not only affects the
PPAs performance but also affects the running time significantly. We can clearly
say that when the number of the slow algorithms' individuals increases in the
population, the running time of the PPA increases. Similarly, if the number
400 of the slow algorithms' individuals decreases, the running time of the PPA
decreases because communication between algorithms requires synchronization.

We also investigate the performance of the PPA for training seven different

architectures of neural networks in a two time series prediction problems. This study represents that portfolio algorithms are able to produce good prediction
405 accuracy.

In the future we would like to study different optimization algorithms which complement each other significantly and apply the proposed parallel models to different real-world problems.

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Table 4: Function error values of the solutions obtained by ABC, DE, PSO and seven PPA instantiations for the benchmark functions

Function	Algorithms									
	ABC	DE	PSO	PPA_{21}	PPA_{22}	PPA_{23}	PPA_{24}	PPA_{25}	PPA_{31}	PPA_{32}
f_1	2.52E-06	3.45E+01	9.35E+09	3.42E-05	4.45E-06	1.46E-04	2.18E+08	2.33E+08	1.52E-03	1.43E-03
f_2	7.12E+01	1.27E-02	7.67E+03	8.90E+01	8.56E+01	9.39E+01	3.43E+03	3.47E+03	9.01E-02	4.13E-01
f_3	1.45E-03	2.21E-02	1.95E+01	4.62E-03	2.50E-03	4.13E-03	1.61E+01	1.63E+01	7.01E-04	3.22E-04
f_4	4.04E+13	8.56E+13	3.02E+12	3.50E+13	3.76E+12	3.09E+12	5.17E+12	3.23E+12	3.60E+12	3.85E+12
f_5	5.66E+08	5.37E+08	3.16E+08	5.33E+08	3.42E+08	3.23E+08	3.52E+08	3.25E+08	3.66E+08	3.41E+08
f_6	2.01E+07	1.97E+07	4.31E+06	2.00E+07	9.54E+06	4.57E+06	7.21E+06	5.40E+06	6.27E+06	5.64E+06
f_7	4.65E+10	6.89E+10	1.49E+10	3.20E+10	6.23E+09	4.09E+09	1.06E+10	1.20E+10	5.15E+09	6.66E+09
f_8	1.58E+07	5.74E+07	7.31E+13	2.65E+07	9.46E+06	9.36E+06	4.50E+11	3.47E+11	1.73E+07	1.75E+07
f_9	7.17E+08	1.74E+09	5.98E+09	6.83E+08	6.61E+08	6.60E+08	9.89E+08	8.69E+08	6.91E+08	6.57E+08
f_{10}	7.27E+03	7.41E+03	8.22E+03	7.28E+03	4.65E+03	4.57E+03	7.10E+03	6.88E+03	4.69E+03	4.57E+03
f_{11}	2.02E+02	2.07E+02	2.12E+02	2.03E+02	1.96E+02	1.95E+02	2.13E+02	2.12E+02	1.97E+02	1.94E+02
f_{12}	7.38E+05	1.32E+06	9.16E+05	7.11E+05	4.97E+05	3.98E+05	3.97E+05	3.64E+05	4.66E+05	3.81E+05
f_{13}	6.32E+02	2.62E+03	8.59E+09	1.00E+03	7.89E+02	1.23E+03	3.24E+07	3.08E+07	1.72E+03	1.51E+03
f_{14}	1.82E+09	5.04E+09	1.67E+09	1.77E+09	1.78E+09	1.71E+09	2.05E+09	1.69E+09	1.68E+09	1.68E+09
f_{15}	1.46E+04	1.58E+04	8.51E+03	1.45E+04	8.73E+03	8.52E+03	8.69E+03	8.53E+03	8.84E+03	8.64E+03
f_{16}	4.05E+02	4.16E+02	3.89E+02	4.05E+02	3.90E+02	3.89E+02	3.90E+02	3.89E+02	3.91E+02	3.89E+02
f_{17}	1.46E+06	2.92E+06	1.12E+06	1.48E+06	1.23E+06	1.10E+06	1.22E+06	1.12E+06	1.27E+06	1.16E+06
f_{18}	7.11E+03	1.00E+04	3.37E+10	1.19E+04	9.55E+03	1.54E+04	2.03E+08	1.92E+08	9.24E+03	9.62E+03
f_{19}	8.54E+06	1.22E+07	2.53E+06	7.32E+06	2.92E+06	2.66E+06	2.85E+06	2.66E+06	3.00E+06	2.83E+06
f_{20}	5.33E+02	3.48E+03	4.31E+10	1.15E+03	8.55E+02	1.54E+03	2.67E+08	2.26E+08	1.89E+03	1.85E+03

Table 5: Average running times (s) of serial PSO and PPA_{31} algorithms for hybrid model

Alg.	CPU(s)	Functions									
		f_1	f_2	f_3	f_4	f_5	f_6	f_7	f_8	f_9	f_{10}
PSO	1	993.62	600.57	530.67	1160.67	817.82	848.53	779.34	758.06	1882.60	1521.97
PPA_{31}	6	226.31	164.55	145.21	251.06	265.34	177.37	171.00	165.26	413.31	421.48
	9	157.60	141.31	102.99	208.13	221.24	144.18	115.74	115.00	306.79	355.04
	12	140.97	122.79	80.68	168.37	202.52	130.99	118.92	118.79	220.08	274.26
	24	135.27	125.93	88.94	148.52	183.87	121.99	127.61	118.77	190.23	229.08
Alg.	CPU(s)	Functions									
		f_{11}	f_{12}	f_{13}	f_{14}	f_{15}	f_{16}	f_{17}	f_{18}	f_{19}	f_{20}
PSO	1	1452.88	767.29	644.27	2641.05	2335.41	2276.08	1036.42	648.41	10093.88	628.80
PPA_{31}	6	313.77	188.24	172.22	596.62	613.06	523.78	284.64	204.70	2357.72	173.97
	9	266.23	171.32	127.80	420.74	485.54	360.24	230.04	141.36	1520.00	147.04
	12	182.86	134.08	127.26	298.20	359.44	266.86	180.34	132.60	1180.98	122.66
	24	170.22	128.08	128.34	215.46	297.28	187.10	158.54	127.00	619.10	123.34

Table 6: The training and testing MSE values obtained by PPA_{31} for the time series

Number of Hidden Nodes	Mackey Glass		Box Jenkins	
	Train	Test	Train	Test
2	3.4945E-03	3.5128E-03	8.9114E-04	9.1565E-04
4	1.3430E-03	1.4180E-03	6.6052E-04	7.9206E-04
6	6.9713E-04	7.7762E-04	6.1556E-04	7.9727E-04
8	5.7153E-04	6.2639E-04	6.0978E-04	7.0342E-04
12	5.4804E-04	6.1113E-04	5.9920E-04	7.2255E-04
16	6.0202E-04	7.0184E-04	5.7387E-04	7.3291E-04
20	7.5745E-04	8.9585E-04	5.8302E-04	9.0047E-04

Table 7: The comparison results of MSE values for the time series

Time Series	Method	BP	PSO	CRPSO	GA	PPA_{31}
Mackey Glass	Train	0.0038	0.0017	5.2504E-04	5.9584E-04	5.4804E-04
	Test	0.0041	0.0018	5.4910E-04	6.2173E-04	6.1113E-04
Box Jenkins	Train	0.0030	0.0029	0.0017	0.0018	6.0978E-04
	Test	0.0056	0.0054	0.0021	0.0023	7.0342E-04

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