A Novel Hybrid Chemical Reaction Optimization Algorithm with Adaptive Differential Evolution Mutation Strategies for Higher Order Neural Network Training

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Abstract: In this paper, an application of a hybrid Chemical Reaction Optimization (CRO) algorithm with adaptive Differential Evolution (DE) mutation strategies for training Higher Order Neural Networks (HONNs), especially the Pi-Sigma Network (PSN) is presented. Contrasting to traditional CRO algorithms, the reactant size (population size) remains fixed throughout all iterations, which makes it easier to implement. In addition, four DE mutation strategies (DE/rand/1, DE/best/1, DE/rand/2 and DE/best/2) with adaptive selection of control parameters as inter-molecular reactions and one intra-molecular reaction have been used. The proposed algorithm combines the diversification property of inter-molecular reactions following DE/rand mutation strategies and intensification property of intra-molecular reaction as well as inter-molecular reactions following DE/best mutation strategies, thereby glorifying the chances of reaching the global optima in less iteration. The performance of the proposed algorithm for HONN training is evaluated through a well-known neural network training benchmark i.e., to classify the parity-p problems. The results obtained from the proposed algorithm to train HONN have been compared with results from the following algorithms: Basic CRO algorithm, CRO-HONN Training (HONNT) and the most popular variants of DE algorithm (DE/rand/1/bin, DE/best/1/bin). It is observed that the application of the proposed hybridized algorithm to (DE-CRO-HONNT) performs statistically better than that of other algorithms considering both classification accuracy and number of generation taken to attain the solutions.

Keywords: CRO, DE, HONN, training algorithm, PSN.

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1. Introduction

Over the past few decades, Artificial Neural Network (ANN) models have been widely used for pattern reorganization, pattern classification and mathematical function approximation. However, now-a-days instead of traditional neural networks, Higher Order Neural Networks (HONNs) have found in increasing consideration in forecasting, classification and regression problems due to several unique characteristics, including: Stronger approximation with faster convergence property; greater storage capacity; and higher fault tolerance capability. On the other hand, the major drawback of most of the HONN models is that the number of weights of the network grows exponentially with the increase in dimensionality of input patterns. But, Pi-Sigma Networks (PSNs) are a special class of HONN which are not only computationally much more efficient than other HONN models but also manages to incorporate the capability of first order HONN indirectly. The PSNs were introduced by Shin and Ghosh [24] and have addressed several difficult tasks such as zeroing polynomials [9] and polynomial factorization [18] more effectively than traditional Feed-forward Neural Networks (FNNs).

Shin and Ghosh [26] have formulated Ridge Polynomial Neural Networks (RPNN) by adding gradually more complex PSNs. RPNNs have shown competitive performance in various tasks such as pattern recognition [29], image prediction [17], time series prediction [10], data classification [26], and intelligent control [12]. Since, the RPNNs are a generalization of PSNs, its effectiveness directly depends on the effectiveness of PSNs. Therefore, a better learning algorithm for PSNs will also improve the efficiency of RPNN. Despite of better performance of PSN and RPNN across various application domains, a few papers were devoted to develop an efficient training algorithm for PSNs [5, 23, 24]. This motivates towards the development of an efficient training algorithm for training PSNs.

The PSNs are supervised networks and efficiency of any supervised neural network depends on the algorithm used for its training. The objective of any supervised training algorithm is to minimize the approximation error by obtaining the optimal weight set. The optimal weight set of PSNs can be obtained by using either gradient or evolutionary learning algorithm. Since the training of PSN is a multimodal search problem, the gradient based training algorithms
often suffer from several shortcomings, including: Easily getting trapped to local minima; have slow convergence properties; and training performance is sensitive to initial values of its parameters. Due to these disadvantages, research on different optimization techniques that are dedicated to PSN training is still needed. There are many optimization techniques such as Differential Evolution (DE) [2, 3, 4, 11, 20, 21, 28], Genetic Algorithm (GA) [7], Particle Swarm Optimization (PSO) [13], Ant Colony Optimization (ACO) [27], a Bee Colony Optimization (BCO) [19], an Evolutionary Strategy (ES) [1], Quantum Inspired Algorithms (QEA) [8], Chemical Reaction Optimization (CRO) [14, 15, 16] etc., which can be used to train PSN. In this paper an attempt has been made to hybridize the self adaptive DE mutation operators with a CRO algorithm to obtain an efficient training algorithm for PSN.

The remainder of this paper is organized as follows: Section 2 briefly describes the mathematical model of PSN, differential evolution algorithm and chemical reaction optimization algorithm. The proposed training algorithm for PSN has been explained in section 3. In section 4 experimental results are presented. And finally, conclusions are drawn in section 5.

2. Related Work

2.1. Pi-Sigma Neural Network

PSN is a special type higher order feed forward neural network that calculates the product of the sum of the input components and passes it to a nonlinear function. The network architecture of PSN as shown in Figure 1 consists of a single hidden layer of summing units and an output layer of product units. The weights connecting the input neurons to the hidden neurons are trainable whereas those connecting the hidden neurons to the output neurons are fixed to one. Such a network topology with only one layer of trainable weights drastically reduces the training time [6, 24, 25]. Moreover, the product units provide the higher order capabilities of HONN models.

Consider a PSN with \( n \) inputs, \( k \) hidden neurons and \( m \) output neurons. The number of hidden neurons in the hidden layer defines the order of a PSN. For a \( k \)th order PSN the number of trainable weights is \( n \times k \) considering each summing unit is associated with \( n \) weights (bias components are not considered). The output of the PSN is computed by making product of the output of \( k \) hidden units and handing it to a nonlinear function, which is defined as:

\[
y = \sigma \left( \prod_{j=1}^{k} h_j \right)
\]

(1)

Where \( \sigma \) is a nonlinear transfer function and \( h_j \) is the output of the \( j \)th hidden unit which is computed by making the sum of the products of each input \( (I_i) \) with the corresponding weight \( (w_{ij}) \) between \( i \)th input and \( j \)th hidden unit. The output of hidden unit is computed as:

\[
h_j = \sum_{i=1}^{n} (w_{ij}I_i)
\]

(2)

2.2. Differential Evolution

The DE algorithm was introduced by Storn and Price [28]. It is a simple yet efficient stochastic direct search method for global optimization of multimodal function. Compared to most other evolutionary algorithms, DE is much simpler and straightforward to implement. Although, PSO is also very easy to code, the performance of DE and its variants outperforms the PSO variants over a wide variety of problems [3, 22] and the Congress on Evolutionary Computation (CEC) competition series. Since, the inception DE, it has been upgraded intensively in recent years [2]. The variants of DE algorithm differ from each other by the type of mutation and crossover scheme being used. The crossover may be binary or exponential. For both the crossover different mutation schemes, suggested by Price et al. [20, 21] are summarized as follows:

- DE/rand/1: \( MV = C_{a1} + F^*(C_{a2} - C_{r1}) \)
- DE/best/1: \( MV = C_{best} + F^*(C_{best} - C_{r1}) \)
- DE/rand/2: \( MV = C_{r1} + F^*(C_{r2} - C_{r3}) + F^*(C_{r4} - C_{r5}) \)
- DE/best/2: \( MV = C_{best} + F^*(C_{r1} - C_{r2}) + F^*(C_{r3} - C_{r4}) \)
- DE/target-to-best/1: \( MV = C_i + F^* (C_{best} - C_i) + F^* (C_{best} - C_{r2}) \)

The conventions used above are DE/a/b, where DE stands for differential evolution, \( a \) represents the base vector to be perturbed (it may be best vector or target vector or a randomly chosen vector), \( b \) represents the number of difference vectors used for perturbation of \( a \); \( MV \) stands for mutant vector; \( C_{best} \) for the best vector of a population and \( C_i \) for a randomly chosen vector from the population. Note that, all the vectors chosen for any mutation strategy must be from the same generation and should be distinct to each other.

2.3. Chemical Reaction Optimization

CRO algorithm was proposed recently by Lam and Li [15], is a metaheuristic optimization technique. It is
inspired by the nature of chemical reactions which loosely couple chemical reactions with optimization. A chemical reactant system consists of a set of chemical substances (reactants/molecules) and its surrounding. Each molecule consists of some atoms and is associated with enthalpy (minimization problem) or entropy (maximization problem). A chemical change of a molecule is triggered by a collision and the corresponding subtle change is called ineffective elementary reaction. There are two types of collision: Uni-molecular/intra-molecular/monomolecular collision (occurs when a molecule hits on some external substance like wall of a container) and inter-molecular collision (occurs when molecules collide with each other). Basing on the number of molecules take part in a reaction, the reaction may be: Uni-molecular or bi-molecular or tri-molecular and so on. Most of the reactions are reversible in nature i.e., they can go in forward or backward direction. Chemical reactions transform one set of chemical substances to another in order to make the system stable. The CRO can be thought of as a new evolutionary technique with molecules as chromosomes; atoms as genes; enthalpy/entropy as fitness function; reactions as crossover and mutation strategies; and reversible reactions as a selection process. However, unlike other evolutionary algorithms in CRO, the reactant size (similar to population size) may vary from one generation to the other. Few authors also have proposed fixed population sized CRO algorithms and shown that fixed population sized CRO not only performs better but also easier to implement [23]. To have an elaborated description regarding CRO algorithm, interested readers may go through the tutorial of CRO [14].

3. DE-CRO-HONNT Method

Algorithm 1 presents the pseudo-code of the proposed method. In this proposed method an attempt has been made to use adaptive DE mutation strategies as inter-molecular reactions of a CRO algorithm and use it for training PSN. Like other evolutionary algorithms, the proposed DE-CRO-HONNT operates in three phases: Initialization phase, iteration phase and final phase. The initial phase assigns the value to initial parameters like termination criteria, total number of reactants/molecules in a generation represented by ReactNum and generates initial set of reactants. The iteration phase simulates the reaction processes. Five different reactions are considered comprising of one intra-molecular (uni-molecular) and four inter-molecular reactions. Every elementary reaction is followed by a greedy reversible reaction to update the reactants.

Algorithm 1: DE-CRO-HONNT.

Set the iteration-counter i=0

/*Randomly Initialize the ReactNum of Reactants from a uniform distribution(upper bound, lower bound): P={R1, R2, R3, . . . , RReactNum}, with Rj={Wj1, Wj2, . . . , Wjn} for j=1, 2, 3, . . . , ReactNum, D=length of each reactant (NOIN=NOHIN), Wj1. . . Wjn, where Wj1 . . . Wjn is kth atom of jth reactant in ith iteration representing a weight of PSN. for j=1 to ReactNum
Calculate the enthalpy e(Rj)
end of for
While (termination criteria is not satisfied) do begin
for j=1 to ReactNum
// Perform reactions over all the reactants of Pj
Generate randj randomly in an interval [0, 1]
if randj ≤ 0.2
Decomposition (Rj); //Uni-molecular Reaction
else if randj >0.2 & & randj ≤0.6
Perform tri-molecular reactions
Generate randj randomly in an interval [0, 1]
if randj ≤ 0.5
//Use DE/rand/1 mutation strategy
Select three random numbers R′1, R′2, R′3 ∈ ReactNum such that R′1≠R′2≠R′3
DErand(R′1, R′2, R′3)
else
// Use DE/best/1 mutation strategy
Select the best reactant Rbest and two random numbers R′1, R′2 ∈ ReactNum such that
R′1 ≠ R′2 ≠ Rbest
DEbest1(R′1, R′2, Rbest)
end of if
else
Perform penta-molecular reactions
Generate randj randomly in an interval [0, 1]
if randj ≤ 0.5
//Use DE/rand/2 mutation strategy
Select five random numbers R1′, R2′, R3′, R4′, R5′ ∈ ReactNum such that
R1′ ≠ R2′ ≠ R3′ ≠ R4′ ≠ R5′
DErand2(R1′, R2′, R3′, R4′, R5′)
else
// Use DE/best/2 mutation strategy
Select the best reactant Rbest and four random numbers R1′, R2′, R3′, R4′ ∈ ReactNum such that
R1′ ≠ R2′ ≠ R3′ ≠ R4′ ≠ Rbest
DEbest2(R1′, R2′, R3′, R4′, Rbest)
end of if
end of if
Apply greedy Reversible Reaction for increased
enthalpy to update reactants
end of for
Set the iteration counter i=i+1
end of while
Use the reactant having best enthalpy as the optimal weight set of PSN.
All the reactions are elaborated in the following subsequent subsections. In the final phase the reactant having best enthalpy is used as the optimal solution (i.e., optimal weight set of a PSN).

3.1. Reactant Encoding

A set of real numbers is used to represent one reactant, with each real number corresponding to a weight of the PSN. Thus, a reactant represents a weight set of the PSN. The length of a reactant depends on the
number of inputs \((n)\) and hidden neurons \((k)\) of the PSN and which is equal to \(n \times k\) (not considering bias units).

### 3.2. Enthalpy of Reactant

Each reactant is associated with some enthalpy (fitness value). As each reactant represents a weight set of the PSN, the Mean Square Error (MSE) on the train set is considered to be its enthalpy. The lower the value of enthalpy represents better the reactant. The MSE is defined as follows:

\[
MSE = \frac{\sum_{i=1}^{NOP} (Y_i - T_i)^2}{NOP}
\]  

(3)

Where \(Y_i\) and \(T_i\) are the output of PSN and target for \(i^{th}\) train pattern.

### 3.3. Elementary Chemical Reactions

One uni-molecular, two tri-molecular and two penta-molecular reactions are considered. The tri-molecular and penta-molecular reactions use different DE mutation strategies. The scale parameter \((F)\) used by the DE mutation strategies are dynamically and self-adaptively determined depending on the problem. Note that the parameter adaptation is somewhat inspired by MDE\_pBX algorithm [11] but the former one is distinct due to its own characteristics. The five reactions are chosen considering both intensification and diversification. All the five reactions considered have equal chance to occur. Therefore, uni-molecular reactions occur with 20%, tri-molecular with 40% and penta-molecular with 40% probability.

#### 3.3.1. Uni-Molecular Reactions

In uni-molecular reactions only one reactant takes part in the reaction and one product is produced by modifying one atom of the reactant. These reactions assist in intensification of the solution by making local search. One uni-molecular reaction is considered called as decomposition reaction which is explained below.

#### 3.3.1.1. Decomposition Reaction

In this reaction a randomly selected atom of the reactant undergoes sudden change to bring a new reactant.

Consider a reactant \(R = \{W_{j,x}, W_{j,2x}, \ldots, W_{j,Dx}\}\) with \(W_{j,x} (x \in [1, n])\) be an atom of the reactant-\(j\). The pseudo-code of the decomposition reaction is described in Algorithm 2.

Algorithm 2: Decomposition(R_1).

**Input**: A reactant \(R_1\).

**Duplicate** \(R_1\) to produce \(R_{new}\).

**Select** an atom \(x (x \in [1, D])\) randomly.

\(W_{new,x} = L + \frac{\lambda}{2} \times (U-L)\).

Where the rate of reaction \((\lambda)\) is a random number generated randomly from a uniform distribution between \([0, 1]\).

**Output**: A new reactant \(R_{new}\).

#### 3.3.2. Tri-Molecular Reactions

In tri-molecular reactions three reactants take part in the reaction to produce one product. Two tri-molecular reactions are considered using two different DE mutation strategies DE/rand/1 and DE/best/1 with the intention to combine the diversification property of the former one and intensification property of later one. Each of the reactions as a whole has a probability of 20% to occur.

##### 3.3.2.1. DErand1 Reaction

Here, DE/rand/1 mutation strategy is used to generate new reactants. In addition, the scale factor \((F)\) used in DE mutation strategy (equivalent to reactant rate \(\lambda\)) is dynamically and self-adaptively determined based on the problem. The rate of reaction \((\lambda)\) is generated randomly from a Cauchy distribution with location parameter \(M\) and scale parameter 0.1. The value of \(M\) is initially set to 0.6 and self-adaptively determined in the following manner:

\[
S_{t} = 0.8 + 0.2 \times \text{rand} (0, 1)
\]

\[
M_{t+1} = S_{t} \times M_{t} + (1 - S_{t}) \times \text{mean} (\lambda_{\text{success}})
\]

(4)

Where \(t\) = Number of times the reaction occurs, \(\lambda_{\text{success}}\) stores the successful rate of reactions that generates better reactants, thereby improving the chances of generating better reaction rates consequently better reactants as more and more this reaction occurs. Here, instead of traditional normal or uniform distribution Cauchy distribution is used because it diversifies the solution more. The pseudo-code of the DErand1 reaction is described in Algorithm 3.

Algorithm 3: DErand1 (R_1, R_2, R_3).

**Input**: Three reactants \(R_1, R_2, R_3\).

\(R_{new} = R_1 + \lambda \times (R_2 - R_3)\).

Where \(\lambda = \text{Cauchy}\text{rand}(M, 0.1)\), is a random number generated from a Cauchy distribution with location parameter \(M\) and scale parameter 0.1. It is regenerated if the random number falls out of the range \([0, 2]\).

**Output**: A new reactant \(R_{new}\).

##### 3.3.2.2. DEbest1 Reaction

This reaction is almost similar to that of DErand1 but, here DE/best/1 mutation strategy is used to generate new reactants. The reaction rate is self-adaptively determined similar to that of previous reaction. The pseudo-code of the DEbest1 reaction is described in Algorithm 4.

Algorithm 4: DEbest1 (R_1, R_2, R_{best}).

**Input**: Three reactants \(R_1, R_2, R_{best}\).

\(R_{new} = R_{best} + \lambda \times (R_1 - R_2)\).

Where \(\lambda = \text{Cauchy}\text{rand} (M, 0.1)\), is a random number generated from a Cauchy distribution with location parameter \(M\) and
scale parameter 0.1. It is regenerated if the random number falls out of the range [0, 2].
Output: A new reactant $R_{new}$.

### 3.3.3. Penta-Molecular Reactions

In penta-molecular reactions five reactants take part in the reaction to produce one product. Two penta- molecular reactions are considered using two other DE mutation strategies DE/rand/2 and DE/best/2 to combine the diversification property of the former one and intensification property of later one.

#### 3.3.3.1. DErand2 Reaction

Here, DE/rand/2 mutation strategy is used to generate new reactants. In addition, the scale factor ($F$) used in DE mutation strategy (equivalent to reactant rate $\lambda$) is dynamically and self adaptively determined based on the problem. The rate of reaction ($\lambda$) is generated randomly from a Gaussian distribution with mean $M$ and standard deviation 0.1. The value of $M$ is initially set to 0.5 and self adaptively determined in the following manner.

$$S_t = 0.9 + 0.1 \times \text{rand}(0, 1)$$

$$M_{t+1} = S_t \times M_t + (1 - S_t) \times \text{mean} \quad (\lambda_{success})$$

Where $t =$ Number of times the reaction occurs, $\lambda_{success}$ stores the successful rate of reactions that generates better reactants, thereby glorifying the chances of generating better reaction rates consequently better reactants as more and more this reaction occurs. Here, instead of Cauchy distribution, Gaussian distribution is used because it generates most of the values within unity due to its short tail property [20]. The pseudo-code of the DErand2 reaction is described in Algorithm 5.

**Algorithm 5:** DErand2 ($R_1, R_2, R_3, R_4, R_5$).

**Input:** Five reactants $R_1, R_2, R_3, R_4, R_5$.

$R_{new} = R_{best} + S_t \times (R_{i} - R_{best}) + \delta \times (R_{j} - R_{k})$.

Where $S_t =$ Gaussian Rand ($M_t, 0.1$), is a random number generated from a Gaussian distribution with mean $M_t$ and standard deviation 0.1. It is regenerated if the random number falls out of the range [0, 1].

**Output:** A new reactant $R_{new}$.

#### 3.3.3.2. DEbest2 Reaction

Here, DE/best/2 mutation strategy is used to generate new reactants. In addition, the scale factor ($F$) used in DE mutation strategy (equivalent to reactant rate $\lambda$) is dynamically and self adaptively determined similar to that of in DErand2 reaction. The pseudo-code of the DEbest2 reaction is described in Algorithm 6.

**Algorithm 6:** DEbest2 ($R_1, R_2, R_3, R_4, R_{best}$).

**Input:** Five reactants $R_1, R_2, R_3, R_4, R_{best}$.

$R_{new} = R_{best} + S_t \times (R_{i} - R_{best}) + \delta \times (R_{j} - R_{k})$.

Where $S_t =$ Gaussian Rand ($M_t, 0.1$), is a random number generated from a Gaussian distribution with mean $M_t$ and standard deviation 0.1. It is regenerated if the random number falls out of the range [0, 1].

**Output:** A new reactant $R_{new}$.

### 3.3.4. Greedy Reversible Reaction

In order to keep the number of reactants fixed throughout all iterations, a greedy reversible reaction between target reactant ($R$) and newly generated reactant ($R_{new}$) is carried out to select the better reactant. By keeping the reactant size fixed it makes the algorithm easier to implement. The pseudo-code of the greedy reversible reaction is elaborated in Algorithm 7.

**Algorithm 7:** Reversible ($R_j, R_{new}$).

**Input:** Two reactants $R_j, R_{new}$.

If enthalpy($R_{new}$) < enthalpy($R_j$)

Set $R_j = R_{new}$

end of if

**Output:** The reactant $R_j$.

### 4. Experimental Results

All simulations were carried out on a system with Intel® core (TM) 2Duo E7500 CPU, 2.93GHz with 2GB RAM and implemented using SCILAB. Parity-$p$ problems ($p \in [3; 7]$) are considered for comparative performance analysis. These problems are widely used and regarded as benchmarks for testing the generalization capability of training algorithms. To classify parity-$p$ ($p \in [3; 7]$) problem, PSNs having structure $p$-$p$-$1$, threshold activation function at output layer and linear transfer function at hidden layer are considered. The population size is fixed to 10 for all the problems and algorithms. The results obtained from proposed method are compared with DE/rand//1/bin, DE/best/1/bin, CRO algorithm used for ANN training [30] and CRO-HONNT [23]. For DE algorithms the crossover probability $C_r$ and scale factor $F$ are fixed to 0.7 and 0.5 respectively.

The termination criterion applied to the training algorithms for parity-$p$ ($p \in [3; 4]$) was the MSE on train set (0.025, 0.0125 respectively); and for parity-$p$ ($p \in [5; 7]$) the networks were trained up to a maximum of 1000 generations or $MSE$ not less than 0.125. The upper and lower bound of initial weight sets for parity-$p$ problem is set to $2^p$ to $-2^p$. By making above experimental set up 1000 independent simulations using each method for each parity-$p$ problem were conducted. To have a better comparison among the methods, Post Hoc analyses were performed on the results obtained from 1000 independent simulations for each problem using each method. Note that in each simulation the initial weight set for all the methods were kept same.
4.1. Performance Measure

To evaluate the effectiveness of the proposed training method with the other methods two performance and percentage of correct classification. The correct classification percentage is computed as follows:

\[
\text{Correct Classification}(\%) = \frac{\sum_{i=1}^{N} Y_i}{NOP} \times 100
\]

Where \( NOP \) is the number of testing patterns (was equal to the training set and each contain \( 2^n \) patterns); \( C_i \) the coefficient representing the correctness of the classification of the \( i^{th} \) testing pattern which is determined as follows:

\[
C_i = \begin{cases} 
1, & \text{when } Y_i = 1 \text{ and } T_i = 1 \\
1, & \text{when } Y_i = -1 \text{ and } T_i = -1 \\
0, & \text{Otherwise} 
\end{cases}
\]

Where \( Y_i \) is the output of PSN and \( T_i \) is the target for \( i^{th} \) test pattern.

4.2. Discussions

One can see from Tables 1 and 2 that all the methods gave perfect generalization (100% correct classification) capability for parity 3 and 4 problems respectively. For both the problems the proposed method takes less number of generations (though statistically insignificant) to obtain the optimal solutions than CRO-HONNT, DE/rand/1/bin and DE/best/1/bin methods whereas takes statistically less number of generations than CRO method.

Table 1. Simulation results on parity 3 problem (best results in bold).

<table>
<thead>
<tr>
<th>Algorithms</th>
<th>Generations</th>
<th>Correct Classification (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± St.D.</td>
<td>Min</td>
</tr>
<tr>
<td>DE-CRO-HONNT</td>
<td>1.72 ± 1.45</td>
<td>1</td>
</tr>
<tr>
<td>CRO-HONNT</td>
<td>1.80 ± 1.64</td>
<td>12</td>
</tr>
<tr>
<td>CRO</td>
<td>2.65 ± 4.03</td>
<td>66</td>
</tr>
<tr>
<td>DE/rand/1</td>
<td>2.12 ± 1.27</td>
<td>17</td>
</tr>
<tr>
<td>DE/best/1</td>
<td>2.11 ± 1.40</td>
<td>9</td>
</tr>
</tbody>
</table>

Mean within a column the same letter(s) are not statistically significant (p=0.05) according to duncan’s multiple range test (SPSS V.16.0.1).

Table 2. Simulation results on parity 4 problem (best results in bold).

<table>
<thead>
<tr>
<th>Algorithms</th>
<th>Generations</th>
<th>Correct Classification (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± St.D.</td>
<td>Min</td>
</tr>
<tr>
<td>DE-CRO-HONNT</td>
<td>16.96 ± 15.92</td>
<td>1</td>
</tr>
<tr>
<td>CRO-HONNT</td>
<td>17.41 ± 15.79</td>
<td>3</td>
</tr>
<tr>
<td>CRO</td>
<td>23.04 ± 40.49</td>
<td>1</td>
</tr>
<tr>
<td>DE/rand/1</td>
<td>18.21 ± 15.38</td>
<td>193</td>
</tr>
<tr>
<td>DE/best/1</td>
<td>18.79 ± 17.54</td>
<td>1</td>
</tr>
</tbody>
</table>

Means within a column the same letter(s) are not statistically significant (p=0.05) according to duncan’s multiple range test (SPSS V.16.0.1).

Table 3. Simulation results on parity 5 problem (best results in bold).

<table>
<thead>
<tr>
<th>Algorithms</th>
<th>Generations</th>
<th>Correct Classification (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± St.D.</td>
<td>Min</td>
</tr>
<tr>
<td>DE-CRO-HONNT</td>
<td>165.83 ± 158.63</td>
<td>5</td>
</tr>
<tr>
<td>CRO-HONNT</td>
<td>173.61 ± 160.95</td>
<td>2</td>
</tr>
<tr>
<td>CRO</td>
<td>194.45 ± 235.14</td>
<td>6</td>
</tr>
<tr>
<td>DE/rand/1</td>
<td>243.30 ± 227.84</td>
<td>10</td>
</tr>
<tr>
<td>DE/best/1</td>
<td>248.62 ± 242.79</td>
<td>5</td>
</tr>
</tbody>
</table>

Means within a column the same letter(s) are not statistically significant (p=0.05) according to duncan’s multiple range test (SPSS V.16.0.1).

Tables 4 and 5 show the experimental results for parity 6 and 7 problems respectively. It is clearly observed that none of the methods gave perfect generalization capability for both problems throughout all 1000 simulations. The proposed method not only provides statistically better generalization capability (correct classification percentage) but also takes statistically significantly less number of generations to attain the solutions than the other methods considered.

To have a better idea regarding the performance (Convergence) of the proposed training algorithm with respect to other four algorithms, a comparative performance was plotted as shown in Figure 2 for parity 7 problem showing MSE error on train set, which again evidenced the superiority of proposed training algorithm (DE-CRO-HONNT).

Table 4. Simulation results on parity 6 problem (best results in bold).

<table>
<thead>
<tr>
<th>Algorithms</th>
<th>Generations</th>
<th>Correct Classification (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± St.D.</td>
<td>Min</td>
</tr>
<tr>
<td>DE-CRO-HONNT</td>
<td>783.89 ± 975.93</td>
<td>28</td>
</tr>
<tr>
<td>CRO</td>
<td>783.97 ± 340.57</td>
<td>25</td>
</tr>
<tr>
<td>DE/rand/1</td>
<td>535.43 ± 332.98</td>
<td>29</td>
</tr>
<tr>
<td>DE/best/1</td>
<td>547.46 ± 338.30</td>
<td>30</td>
</tr>
</tbody>
</table>

Means within a column the same letter(s) are not statistically significant (p=0.05) according to duncan’s multiple range test (SPSS V.16.0.1).

Table 5. Simulation results on parity 7 problem (best results in bold).

<table>
<thead>
<tr>
<th>Algorithms</th>
<th>Generations</th>
<th>Correct Classification (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± St.D.</td>
<td>Min</td>
</tr>
<tr>
<td>DE-CRO-HONNT</td>
<td>875.02 ± 281.45</td>
<td>35</td>
</tr>
<tr>
<td>CRO</td>
<td>991.48 ± 64.87</td>
<td>203</td>
</tr>
<tr>
<td>DE/rand/1</td>
<td>995.42 ± 36.47</td>
<td>620</td>
</tr>
<tr>
<td>DE/best/1</td>
<td>714.35 ± 300.42</td>
<td>150</td>
</tr>
<tr>
<td></td>
<td>707.16 ± 357.50</td>
<td>152</td>
</tr>
</tbody>
</table>

Means within a column the same letter(s) are not statistically significant (p=0.05) according to duncan’s multiple range test (SPSS V.16.0.1).
Figure 2. DE-CRO-HONNT, CRO-HONNT, CRO, DE/rand/1/bin and DE/best/1/bin algorithms MSE value on train set of parity 7 problem up to termination criteria.

5. Conclusions
In this paper, a hybrid DE-CRO-HONNT training algorithm for PSN is developed. In this algorithm adaptive DE mutation strategies are hybridized as inter-molecular reactions in CRO algorithm. The proposed algorithm enjoys both the intensification property of DE/best mutation strategies along with uni-molecular reaction and diversification property of DE/rand mutation strategies. Thus, a trade between intensification and diversification is implicitly maintained. In addition, control parameters for DE mutation strategies are dynamically and self adaptively determined based on the problem in hand. The simulation results demonstrate that the proposed training algorithm has superior performance in terms of correct classification percentage and generations taken to attain the solutions when compared with most popular DE variants, traditional CRO method and CRO-HONNT method. It is also observed that the performance of other methods drops off sharply with the increase in number of parity bits whereas the same is not the case for DE-CRO-HONNT. Therefore, it can be concluded that, the use of DE-CRO-HONNT method incorporates efficient and effective searching mechanisms, such that it has less chance to trap to local minima and thus enhance the higher order neural network training procedure.

References
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