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Article

Introducing a new genetic operator based on Differential Evolution for effective training of neural networks

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Abstract: Artificial neural networks are widely established models used in a variety of real - world problems derived from physics, chemistry etc. These machine learning models contain a series of parameters that must be appropriately tuned by various optimization techniques in order to be effective in the problems they face. Genetic algorithms have been used in many cases in the recent literature to train artificial neural networks and various modifications have been introduced to enhance this procedure. In this article, the incorporation of a novel genetic operator in genetic algorithms is proposed in order to effectively train artificial neural networks. The new operator is based on the differential evolution technique and it is periodically applied to randomly selected chromosomes from the genetic population. Furthermore, to find a promising range of values for the parameters of the artificial neural network, an additional genetic algorithm is executed before the execution of the basic algorithm. The modified genetic algorithm was used to train neural networks for classification and regression datasets and the results are reported and compared against other methods that train neural networks.

Keywords: Neural networks; Genetic algorithms; Evolutionary computation

1. Introduction

A machine learning model that has been widely used in recent decades in dozens of problems are artificial neural networks [1,2], which are parametric models commonly defined as $N(\overrightarrow{x}, \overrightarrow{w})$. The vector \overrightarrow{x} stands for the input pattern and the vector \overrightarrow{w} represents the associated set of parameters that should be calculated by any optimization method. The calculation is performed by minimizing the so - called training error, expressed as:

 $E(N(\overrightarrow{x}, \overrightarrow{w})) = \sum_{i=1}^{M} (N(\overrightarrow{x}_i, \overrightarrow{w}) - y_i)^2$ (1)

The values $(\overrightarrow{x_i}, y_i)$, i = 1, ..., M form the training set of the problem, where y_i represent the expected outputs for each pattern $\overrightarrow{x_i}$.

Artificial neural networks have applied on a wide series of problems from various areas, such as physics [3,4], astronomy [5], chemistry [6], economics [7], medicine [8,9] etc. The equation 1 has been minimized by various methods in the relevant literature. Among them one can find the Back propagation method [10,11], the RPROP method [12–14], Quasi Newton methods [15,16], Simulated Annealing [17], Particle Swarm Optimization (PSO) [18, 19], Genetic Algorithms [20,21], Differential Evolution [22], Ant Colony Optimization [23], Gray Wolf Optimizer [24], Whale optimization [25] etc. Moreover, Zhang et al proposed a hybrid algorithm that conjuncts PSO and the Back Propagation algorithm for neural networks training [26]. Also, recently many researchers proposed methods that take advantage of parallel processing units in order to speed up the training process [27,28].

Citation: Tsoulos, I.G.; Charilogis, V.; Tsalikakis D. Introducing a new genetic operator based on Differential Evolution for effective training of neural networks. *Journal Not Specified* 2023, 1, 0. https://doi.org/

Received:

Revised:

Accepted:

Published:

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Furthermore, a series of papers has been published recently that tackle the initialization procedure for the parameters of neural networks. These methods include decision trees [29], incorporation of the Cauchy's inequality [30], discriminant learning [31], usage of polynomial bases [32], usage of intervals [33] etc. A systematic review of initialization methods can be found in the work of Narkhede et al [34].

Additionally, finding the optimal architecture of an artificial neural network can effectively contribute to its training, since on the one hand it will reduce the required training time and on the other hand it will eliminate the problem of overfitting. In this direction, a series of researchers have proposed many methods to tackle this problem, such as genetic algorithms [35,36], the application of the PSO method method [37], application of reinforcement learning [38] etc. Also, Tsoulos et al. proposed the usage of Grammatical Evolution technique [39] to construct artificial neural networks [40].

This paper proposes the usage of a two-stage technique for the efficient training of artificial neural networks. In the first stage, a genetic algorithm is used to efficiently identify a range of values within which the parameters of the artificial neural network should be optimized. In the second stage, a genetic algorithm optimizes these parameters, which uses a new operator to enhance its results. This new operator is based on the differential evolution technique [41] and is applied periodically to randomly selected chromosomes of the genetic population. The first stage of the technique is necessary to ensure that the parameters of the artificial neural network will be trained within a range of values, which will prevent their overfitting as much as possible. In the second stage, the differential evolution method was selected as the base of the new operator. This method is an evolutionary technique widely used in a series of practical problems, such as community detection [42], structure prediction [43], motor fault diagnosis [44], and clustering techniques [45]. Furthermore, this method was chosen as the basis for the new genetic operator due to the small number of required parameters that the user must specify.

The remaining of this article is organized as follows: in section 2 the proposed method is discussed in detail, in section 3 the used datasets as well as the conducted experiments are discussed and finally, in section 4 some conclusions are presented.

2. Method description

The two phases of the proposed method are analyzed in detail in this section. During the first phase a genetic algorithm is utilized in order to detect a promising interval of values for the parameters of neural network. In the second phase a genetic algorithm that incorporates the suggested operator is applied to minimize the training error of the neural network and the parameters are initialized inside the interval located during the first phase.

2.1. The first phase of the proposed method

In the first phase of the proposed technique, a genetic algorithm is used to identify a range of values for the parameters of the artificial neural network. Genetic algorithms are evolutionary methods, where a series of randomly created candidate solutions, those called chromosomes, are evolved ireratively through a series of steps similar to natural processes such as selection, crossover and mutation. Genetic algorithms have been used successfully in a series of real - world problems, such as placement of wind turbines [46], water distribution [47], economics [48], neural network training [49] etc. The neural networks adopted in this manuscript have the following form, as proposed in [40]:

$$N(\overrightarrow{x}, \overrightarrow{w}) = \sum_{i=1}^{H} w_{(d+2)i-(d+1)} \sigma \left(\sum_{j=1}^{d} x_j w_{(d+2)i-(d+1)+j} + w_{(d+2)i} \right)$$
(2)

Where the value H denotes the total number of processing units in this network and the value d defines the number of inputs for the pattern \overrightarrow{x} . Hence, the total number of parameters for this network are: n = (d+2)H. The function $\sigma(x)$ is the sigmoid function, defined as:

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$$\sigma(x) = \frac{1}{1 + \exp(-x)} \tag{3}$$

The steps of the algorithm of the first phase have as follows:

1. Initialization step.

- (a) **Set** the number of chromosomes N_c and the maximum number of allowed generations N_g .
- (b) **Set** the selection rate p_s and the mutation rate p_m .
- (c) **Set** the margin factor a, where $a \ge 1$.
- (d) **Set** k = 0 as the generation counter.
- (e) **Initialize** randomly the chromosomes g_i , $i = 1, ..., N_c$. Each chromosome is a vector of parameters for the artificial neural network.

2. Fitness calculation step.

- (a) **For** $i = 1, ..., N_c$ **do**
 - Create the neural network $N_i(\overrightarrow{x}, \overrightarrow{g_i})$ for the chromosome g_i .
 - ii. **Calculate** the associated fitness value f_i as

$$f_i = \sum_{i=1}^{M} (N_i(\overrightarrow{x_j}, \overrightarrow{g_i}) - y_j)^2$$

for the pairs $(\overrightarrow{x_i}, y_i)$, j = 1, ..., M of the training set.

- (b) End For
- 3. Genetic operations step.
 - (a) **Transfer** the best $(1 p_s) \times N_c$ chromosomes of the current generation to the next one. The remaining will be replaced by chromosomes produced in crossover and mutation.
 - (b) **Perform** the crossover procedure. During this procedure, for each pair of constructed chromosomes $(\widetilde{z},\widetilde{w})$ two chromosomes will be selected from the current population using tournament selection. The production of the new chromosomes is performed using process suggested by Kaelo et al [50].
 - (c) **Perform** the mutation procedure. During the mutation procedure, for each element of each chromosome a random number $r \in [0,1]$ is selected. The corresponding element is altered randomly when $r \leq p_m$.

4. Termination check step.

- (a) **Set** k = k + 1
- (b) If $k \le N_g$ then goto Fitness Calculation step.

5. Margin creation step.

- (a) **Obtain** the best chromosome g^* with the lowest fitness value.
- (b) **Create** the vectors L^* and R^* as:

$$L_i^* = -a|g_i^*|, i = 1,...,n$$

 $R_i^* = a|g_i^*|, i = 1,...,n$

2.2. The second phase of the proposed method

During the second phase a second genetic algorithm is used to minimize the training error of the neural network. The parameters of the neural network are initialized inside the vectors L^* and R^* produced in the previous phase of the algorithm. Also, a novel stochastic genetic operator, which is based on the Differential Evolution approach, is applied periodically to the genetic population. This new stochastic operator is used to improve the performance of randomly selected chromosomes and to speed up the overall

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genetic algorithm in finding the global minimum. The main steps of the algorithm executed on the second phase have as follows:

1. Initialization step.

- (a) **Set** the number of chromosomes N_c and the maximum number of allowed generations N_g .
- (b) **Set** the selection rate $p_s \le 1$ and the mutation rate $p_m \le 1$.
- (c) **Set** the crossover probability CR, used in the new genetic operator.
- (d) **Set** the differential weight *F* that will be used in the novel genetic operator.
- (e) **Set** as N_i the number of generations before the application of the new operator.
- (f) **Set** as N_l the number of chromosomes that will participate in the new operator.
- (g) **Initialize** the g_i , $i = 1, ..., N_c$ chromosomes inside the vectors L^* and R^* of the previous phase.
- (h) **Set** k = 0 the generation counter.

2. Fitness calculation step.

- (a) **For** $i = 1, ..., N_c$ **do**
 - i. **Produce** the corresponding neural network $N_i(\overrightarrow{x}, \overrightarrow{g_i})$ for the chromosome g_i .
 - ii. Calculate the fitness value f_i as

$$f_i = \sum_{j=1}^{M} (N_i(\overrightarrow{x_j}, \overrightarrow{g_i}) - y_j)^2$$

(b) End For

3. Application of genetic operators.

- (a) **Copy** the best $(1 p_s) \times N_c$ chromosomes with the lowest fitness values to the next generation. The remaining will be replaced by chromosomes produced in crossover and mutation.
- (b) **Apply** the same crossover procedure as in the algorithm of the first phase.
- (c) **Apply** the same mutation procedure as in the genetic algorithm of the first phase.

4. Application of the novel genetic operator.

- (a) If $k \mod N_i = 0$ then
 - i. Create the set $C = \{z_1, z_2, \dots, z_{N_{cr}}\}$ of N_l randomly selected chromosomes
 - ii. **For** $i = 1, ..., N_l$ apply the deOperator of algorithm 1 to every chromosome $z_i \in C$.
- (b) End if

5. Termination check step.

- (a) **Set** k = k + 1
- (b) If $k \le N_g$ goto Fitness calculation step.

6. **Testing step**.

- (a) **Obtain** the best chromosome g^* from the genetic population.
- (b) **Create** the corresponding neural network $N^*(\overrightarrow{x_j}, \overrightarrow{g^*})$.
- (c) **Apply** this neural network to the test set of the objective problem and report the error.

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Algorithm 1 The proposed genetic operator.

Function deOperator(g, F, CR)

- 1. **Select** three distinct chromosomes *a*, *b*, *c* from the current population using tournament selection.
- 2. **Set** $R \in [1, n]$ a randomly selected integer.
- 3. **Set** t = g, as the trial chromosome.
- 4. **For** i = 1, ..., n **do**
 - (a) **Select** $r \in [0, 1]$ a random number.
 - (b) If i = R or $r \le CR$ then $t_i = a_i + F \times (b_i c_i)$
 - (c) Set $t_f = \sum_{j=1}^{M} \left(N_i \left(\overrightarrow{x_j}, \overrightarrow{t} \right) y_j \right)^2$
 - (d) If $t_f \leq f_g$ then g = t.
- 5. End For
- 6. **Return** g.

End Function

3. Experiments

To demonstrate the dynamics and reliability of the proposed methodology, a series of experiments were carried out on known datasets from the relevant literature. These datasets were obtained from the following databases:

- 1. The UCI database https://archive.ics.uci.edu/(accessed on 5 March 2025)[51]
- 2. The Keel website, https://sci2s.ugr.es/keel/datasets.php(accessed on 5 March 2025)[52]. 166
- 3. The Statlib URL ftp://lib.stat.cmu.edu/datasets/index.html(accessed on 5 March 2025).

3.1. Experimental datasets

The following series of classification datasets were used in the conducted experiments:

- 1. The Alcohol dataset, which is related to experiments on alcohol consumption [53].
- 2. The Appendicitis dataset, which is a medical dataset [54].
- 3. The Australian dataset, which is used in bank transactions [55].
- 4. The Balance dataset, which contains measurements from various psychological experiments [56].
- 5. The Circular dataset, which was created artificially.
- 6. The Cleveland dataset, which is a medical dataset [57,58].
- 7. The Dermatology dataset, which is a medical dataset regarding dermatology problems [59].
- 8. The Ecoli dataset, which is used in protein problems [60].
- 9. The Fert dataset, related to the detection of relations between sperm concentration and demographic data.
- 10. The Haberman dataset, which is related to the detection of breast cancer.
- 11. The Hayes roth dataset [61].
- 12. The Heart dataset, which is related to some heart diseases [62].
- 13. The HouseVotes dataset, related to data from Congressional voting in USA [63].
- 14. The Ionosphere dataset, that contains measurements from the ionosphere [64,65].
- 15. The Liverdisorder dataset, which is a medical dataset [66,67].
- 16. The Lymography dataset [68].
- 17. The Mammographic dataset, which is a medical dataset [69].
- 18. The Parkinsons dataset, that was used in the detection of Parkinson's disease [70,71].
- 19. The Pima dataset, a medical dataset related to the detection of diabetes's disease [72].
- 20. The Popfailures dataset, related to climate model simulations [73].
- 21. The Regions2 dataset, related to some diseases in liver [74].
- 22. The Saheart dataset, related to some heart diseases [75].
- 23. The Segment dataset, related to image processing [76].

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- 24. The Sonar dataset, used to discriminate sonar signals [77].
- 25. The Spiral dataset, which was created artificially.
- 26. The StatHeart dataset, a medical dataset regarding heart diseases.
- 27. The Student dataset, which is related to experiments conducted in schools [78].
- 28. The WDBC dataset, which is related to the detection of cancer [79].
- 29. The Wine dataset, used to detection of the quality of wines [80,81].
- 30. The EEG dataset, which contains various EEG measurements [82,83]. From this dataset the following cases were utilized: Z_F_S, ZO_NF_S and ZONF_S.
- 31. The ZOO dataset, which is used for animal classification [84].

Also, the following regression datasets were incorporated in the conducted experiments:

- 1. The Abalone dataset, that was used to predict the age of abalones [85].
- 2. The Airfoil dataset, derived from NASA [86].
- 3. the Baseball dataset, used to predict the salary of baseball players.
- 4. The BK dataset, related to basketball games [87].
- 5. The BL dataset, related to some electricity experiments.
- 6. The Concrete dataset, which is related to civil engineering [88].
- 7. The Dee dataset, which is related to the price of electricity.
- 8. The Housing dataset, related to the price of houses [89].
- 9. The Friedman dataset, used in various benchmarks [90].
- 10. The FY dataset, related to fruit flies.
- 11. The HO dataset, obtained from the STATLIB repository.
- 12. The Laser dataset, related to laser experiments.
- 13. The LW dataset, related to the prediction of the weight of babes.
- 14. The MB dataset, which was obtained from Smoothing Methods in Statistics.
- 15. The Mortgage dataset, which is an economic dataset.
- 16. The Plastic dataset, related to the pressure in plastics.
- 17. The PY dataset [91].
- 18. The PL dataset, obtained from the STATLIB repository.
- 19. The Quake dataset, used to detect the strength of earthquakes.
- 20. The SN dataset, which is related to trellising and pruning.
- 21. The Stock dataset, used to estimate the price of stocks.
- 22. The Treasury dataset, which is an economic dataset.
- 23. The VE dataset, obtained from the STATLIB repository.

3.2. Experimental results

The code used in the conducted experiments was written in ANSI C++ and all runs were performed 30 times using a different seed for the random number generator each time. The validation of the experiments were done using the well - known method of 10 fold cross validation. For the case of classification datasets the average classification error is reported in the experimental tables. This error is calculated through the following equation:

$$E_C(N(w,x)) = 100 \times \frac{\sum_{i=1}^{K} (class(N(w,x_i)) - y_i)}{K}$$
 (4)

where the set $T = \{x_i, y_i\}$, i = 1, ..., K denotes the test set of the objective problem. For regression datasets the average regression error as calculated on the test set is reported and it is denoted as:Also, the regression error is defined as:

$$E_R(N(w,x)) = \frac{\sum_{i=1}^K (N(w,x_i) - y_i)^2}{K}$$
 (5)

The values for the parameters of the proposed method are shown in Table 1. In the experimental tables the following notation is used:

1. The column DATASET represents the objective problem.

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- 2. The column ADAM denotes the incorporation of the ADAM optimizer [92] to train a neural network with H=10 processing nodes.
- 3. The column BFGS represents the application of the BFGS optimizer [93] to train a neural network with H = 10 processing nodes.
- 4. The column GENETIC denotes the usage of a Genetic Algorithm with the same set of parameters as shown in Table 1 to train an artificial neural network with H=10 processing nodes.
- 5. The column NEAT is used for the application of the NEAT method (NeuroEvolution of Augmenting Topologies) [94].
- 6. The row AVERAGE is used for the average classification or regression error for all datasets.

Table 1. The values of the experimental parameters.

PARAMETER	MEANING	VALUE
N_g	Number of generations allowed	200
N_c	Number of chromosomes	500
N_i	Number of generations before the application of the operator	20
N_l	Number of chromosomes where the operator will be applied	20
F	Differential Weight	0.8
CR	Crossover probability	0.9
p_s	Selection rate	0.1
p_m	Mutation rate	0.05
Н	Number of processing nodes	10
а	Margin factor	1.0

The experimental results for the classification datasets are shown in Table 2 and for the regression datasets in Table 3.

Table 2. Experiments for classification datasets

DATASET	ADAM	BFGS	NEAT	GENETIC	PROPOSED
Alcohol	57.78%	41.50%	66.80%	39.57%	24.79%
Appendicitis	16.50%	18.00%	17.20%	18.10%	15.97%
Australian	35.65%	38.13%	31.98%	32.21%	31.76%
Balance	7.87%	8.64%	23.14%	8.97%	8.39%
Circular	19.95%	6.08%	35.18%	5.99%	3.69%
Cleveland	67.55%	77.55%	53.44%	51.60%	48.10%
Dermatology	26.14%	52.92%	32.43%	30.58%	7.74%
Ecoli	64.43%	69.52%	43.44%	54.67%	47.62%
Fert	23.98%	23.20%	15.37%	28.50%	22.00%
Haberman	29.00%	29.34%	24.04%	28.66%	25.99%
Hayes Roth	59.70%	37.33%	50.15%	56.18%	37.00%
Heart	38.53%	39.44%	39.27%	28.34%	24.79%
HouseVotes	7.48%	7.13%	10.89%	6.62%	5.22%
Ionosphere	16.64%	15.29%	19.67%	15.14%	9.56%
Liverdisorder	41.53%	42.59%	30.67%	31.11%	31.08%
Lymography	29.26%	35.43%	33.70%	23.26%	28.60%
Mammographic	46.25%	17.24%	22.85%	19.88%	16.98%
Parkinsons	24.06%	27.58%	18.56%	18.05%	18.02%
Pima	34.85%	35.59%	34.51%	32.19%	30.44%
Popfailures	5.18%	5.24%	7.05%	5.94%	4.29%
Regions2	29.85%	36.28%	33.23%	29.39%	26.43%
Saheart	34.04%	37.48%	34.51%	34.86%	32.60%
Segment	49.75%	68.97%	66.72%	57.72%	30.00%
Sonar	30.33%	25.85%	34.10%	22.40%	18.78%
Spiral	47.67%	47.99%	48.66%	48.66%	44.20%
Statheart	44.04%	39.65%	44.36%	27.25%	22.72%
Student	5.13%	7.14%	10.20%	5.61%	4.16%
Wdbc	35.35%	29.91%	12.88%	8.56%	7.73%
Wine	29.40%	59.71%	25.43%	19.20%	8.55%
Z_F_S	47.81%	39.37%	38.41%	10.73%	6.46%
ZO_NF_S	47.43%	43.04%	43.75%	21.54%	6.01%
ZONF_S	11.99%	15.62%	5.44%	2.60%	1.79%
ZOO	14.13%	10.70%	20.27%	16.67%	9.07%
AVERAGE	32.70%	33.01%	31.16%	25.48%	20.02%

GENETIC DATASET ADAM **BFGS NEAT** PROPOSED **ABALONE** 4.30 5.69 7.17 9.88 4.33 AIRFOIL 0.005 0.003 0.003 0.067 0.003 BASEBALL 77.90 119.63 67.45 103.60 100.39 BK 0.03 0.28 0.027 0.15 0.02 2.55 BL0.28 5.74 0.05 0.002 0.078 0.066 0.0099 0.003 CONCRETE 0.081 DEE 2.36 1.013 1.512 0.20 0.63 HOUSING 97.38 80.20 43.26 56.49 26.62 FRIEDMAN 22.90 1.263 1.249 19.35 1.33 0.038 0.22 0.65 0.08 0.039 FY НО 0.62 0.035 2.78 0.169 0.014 LASER 0.03 0.015 0.59 0.084 0.0027 LW 0.028 2.98 1.90 0.17 0.016 MB 0.06 0.129 3.39 0.061 0.048 **MORTGAGE** 9.24 8.23 2.41 0.31 14.11 **PLASTIC** 11.71 20.32 2.79 20.77 2.20 0.29 0.28 PL 0.117 0.098 0.023 PY 0.09 0.578 0.075 105.41 0.016 **QUAKE** 0.06 0.42 0.040 0.298 0.043 0.40 2.95 SN 0.206 0.174 0.024 **STOCK** 302.43 180.89 3.88 215.82 3.47 TREASURY 9.91 2.929 11.16 15.52 0.44 0.359 1.92 2.43 VE 0.0450.023 **AVERAGE** 17.41 25.12 12.80 19.80 4.64

Table 3. Experiments for regression datasets.

Also, in Figures 1 and 2 the statistical comparison for the experimental results is outlined graphically.

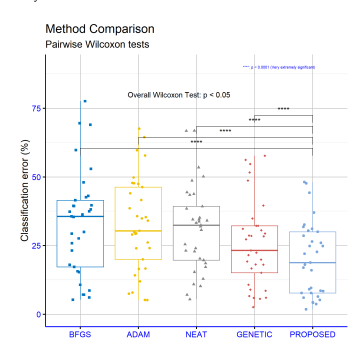


Figure 1. Statistical comparison for the obtained experimental results in the classification datasets.

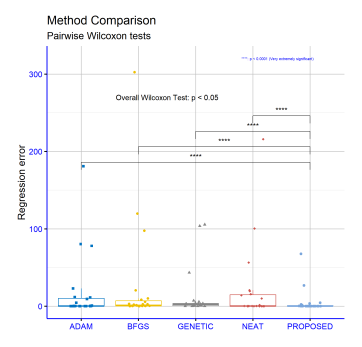


Figure 2. Statistical comparison for the experimental results on the regression datasets.

In the scientific analysis of the experimental results, the proposed model (PROPOSED) demonstrates statistically significant superiority over other methods (ADAM, BFGS, GENETIC, NEAT, RBF) in both classification and regression datasets. For classification, the PROPOSED model achieves a mean error rate of 20.02%, compared to 25.48%-33.01% for the other methods, with extremely low p-values (e.g., $p=2.3\times10^{-10}$ against BFGS). In regression, the model's mean absolute error (4.64) is significantly lower than that of the comparative methods (12.8-25.12), with strong statistical significance ($p<10^{-5}$). However, in certain datasets (e.g., CLEVELAND, FRIEDMAN), the model's performance decreases, indicating dependence on data characteristics.

In the analysis of the experiments (Figure 1) for classification datasets, the proposed model (PROPOSED) demonstrates statistically significant superiority over all comparative methods (BFGS, ADAM, NEAT, GENETIC), with extremely low p-values ($p=2.3\times10^{-10}$ to $p=4.4\times10^{-6}$). This indicates strong differences at a confidence level >99.9%, particularly against BFGS and ADAM ($p<10^{-9}$). For regression datasets (Figure 2), PROPOSED maintains significant superiority over all methods, though with slightly higher p-values ($p=10^{-5}$ to $p=2.4\times10^{-7}$). The largest difference is observed against NEAT ($p=2.4\times10^{-7}$), while the smallest is against GENETIC ($p=8.6\times10^{-5}$).

3.3. Experiments with the differential weight

An additional experiment was performed to demonstrate the reliability of the proposed methodology. In this experiment, three different differential weight F calculation techniques were used for the proposed operator. These techniques are the following:

- 1. Fixed, which is the default technique in the proposed method. In this technique the value F = 0.8 is used for the differential weight.
- 2. Adaptive, where the adaptive calculation of parameter *F* is uses as proposed in [95].
- 3. Random, where the stochastic calculation of parameter *F* as proposed in [96] is used.

In table 4 the results from the application of the proposed method using the previously mentioned techniques for the differential weight are depicted for the classification datasets. Similarly, the same method is applied on the regression datasets and the results are presented in Table 5.

Table 4. Experiments for classification datasets using a series of differential weight mechanisms.

DATASET	FIXED	ADAPTIVE	RANDOM	
Alcohol	24.79%	25.65%	23.02%	
Appendicitis	15.97%	15.83%	15.80%	
Australian	31.76%	31.61%	31.60%	
Balance	8.39%	8.45%	8.65%	
Circular	3.69%	3.67%	3.78%	
Cleveland	48.10%	48.39%	48.35%	
Dermatology	7.74%	7.27%	7.37%	
Ecoli	47.62%	48.34%	47.96%	
Fert	22.00%	22.17%	22.10%	
Haberman	25.99%	26.20%	26.12%	
Hayes Roth	37.00%	38.38%	37.65%	
Heart	24.79%	24.15%	25.51%	
HouseVotes	5.22%	5.21%	4.78%	
Ionosphere	9.56%	9.28%	9.30%	
Liverdisorder	31.08%	31.46%	31.11%	
Lymography	28.60%	28.95%	27.88%	
Mammographic	16.98%	17.02%	17.18%	
Parkinsons	18.02%	17.86%	17.90%	
Pima	30.44%	31.12%	30.48%	
Popfailures	4.29%	4.25%	4.28%	
Regions2	26.43%	25.94%	26.35%	
Saheart	32.60%	33.11%	32.92%	
Segment	30.00%	28.83%	30.85%	
Sonar	18.78%	18.08%	18.70%	
Spiral	44.20%	44.21%	44.12%	
Statheart	22.72%	22.72%	23.41%	
Student	4.16%	3.95%	4.35%	
Wdbc	7.73%	7.48%	7.40%	
Wine	8.55%	6.29%	6.74%	
Z_F_S	6.46%	6.92%	6.65%	
ZO_NF_S	6.01%	6.10%	5.89%	
ZONF_S	1.79%	1.71%	1.76%	
ZOO	9.07%	6.57%	8.90%	
AVERAGE	20.02%	19.91%	19.97%	

DATASET FIXED ADAPTIVE RANDOM ABALONE 4.33 4.24 4.34 AIRFOIL 0.003 0.003 0.003 BASEBALL 67.45 67.23 66.76 BK 0.02 0.02 0.02 BL 0.002 0.002 0.002 **CONCRETE** 0.003 0.003 0.003 DEE 0.20 0.20 0.20 HOUSING 26.62 26.07 26.11 FRIEDMAN 1.33 1.21 1.34 0.039 0.039 0.039 FY НО 0.014 0.014 0.014 LASER 0.0027 0.0027 0.0028 LW 0.016 0.011 0.011 MB 0.048 0.048 0.048 MORTGAGE 0.31 0.35 0.34 PLASTIC 2.20 2.13 2.20 PL 0.023 0.022 0.022 PY 0.016 0.017 0.017 **QUAKE** 0.043 0.082 0.04 0.024 0.024 0.024 SN STOCK 3.33 3.47 3.45 TREASURY 0.44 0.42 0.45 0.023 VE 0.023 0.023 **AVERAGE** 4.64 4.59 4.59

Table 5. Experiments for regression datasets using a variety of weight mechanism methods.

Furthermore, the statistical comparison for these experimental tables is outlined in Figures 3 and 4 respectively.

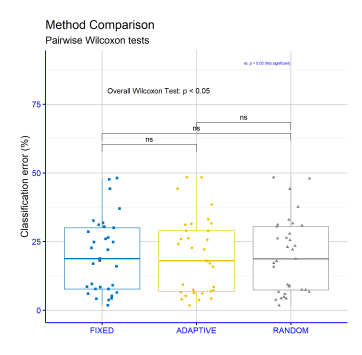


Figure 3. Statistical comparison for the experimental results on the classification datasets, using the proposed method and a series of differential weight techniques.

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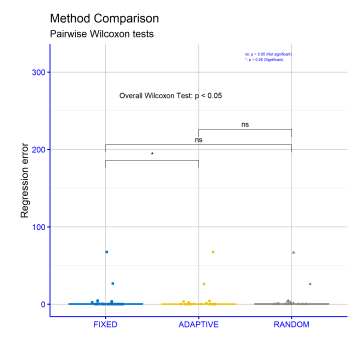


Figure 4. Statistical comparison for the experiments on the regression datasets using the proposed method and a series of differential weight calculation techniques.

When comparing differential weight calculation methods (FIXED, ADAPTIVE, RANDOM), the ADAPTIVE method shows slightly better average performance (19.91% in classification, 4.59 in regression) compared to FIXED (20.02%, 4.64) and RANDOM (19.97%, 4.59). However, the differences are minimal and often statistically insignificant (e.g., p=0.83 for FIXED vs ADAPTIVE in classification). In some datasets, such as Lymography and HouseVotes, the RANDOM method outperforms others, highlighting the need to tailor the method to the specific problem. In regression, the only significant difference occurs between FIXED and ADAPTIVE (p=0.041).

For this experiment as shown in Figure 3, the differences between FIXED and ADAP-TIVE (p=0.83) and ADAPTIVE vs RANDOM (p=0.94) are not statistically significant, suggesting equivalent performance. However, the comparison FIXED vs RANDOM ($p=2.3\times10^{-10}$) reveals a highly significant difference, likely due to the random nature of RANDOM. For regression datasets (Figure 4), the only significant difference occurs between FIXED and ADAPTIVE (p=0.041), while the remaining comparisons (FIXED vs RANDOM: p=0.75, ADAPTIVE vs RANDOM: p=0.31) lack statistical significance.

3.4. Experiments with the margin factor a

An additional experiment was executed to outline the performance and the stability of the proposed method. In this experiment that margin factor denoted as a in the proposed method was varied from a=1.0 (which is the default value) to a=4.0. The experimental results for the classification datasets are shown in Table 6 and for regression datasets in Table 7.

Table 6. Experiments for classification datasets using a series of values for the margin factor a.

DATASET	a = 1.0	a = 1.5	a = 2.0	a = 4.0
Alcohol	24.79%	22.88%	25.55%	28.42%
Appendicitis	15.97%	16.33%	18.07%	19.51%
Australian	31.76%	32.79%	33.23%	33.68%
Balance	8.39%	8.50%	8.39%	8.78%
Circular	3.69%	4.19%	4.22%	4.70%
Cleveland	48.10%	47.17%	46.68%	47.74%
Dermatology	7.74%	7.47%	7.60%	7.83%
Ecoli	47.62%	48.69%	48.29%	49.39%
Fert	22.00%	23.47%	23.50%	24.27%
Haberman	25.99%	26.26%	26.74%	27.14%
Hayes Roth	37.00%	39.15%	39.69%	40.05%
Heart	24.79%	24.64%	24.85%	24.89%
HouseVotes	5.22%	4.91%	5.05%	5.02%
Ionosphere	9.56%	9.65%	9.92%	10.23%
Liverdisorder	31.08%	31.94%	31.42%	32.13%
Lymography	28.60%	28.24%	28.79%	26.69%
Mammographic	16.98%	16.34%	16.35%	16.69%
Parkinsons	18.02%	18.88%	18.56%	19.40%
Pima	30.44%	30.89%	31.11%	31.03%
Popfailures	4.29%	5.07%	5.53%	5.97%
Regions2	26.43%	26.53%	26.30%	25.35%
Saheart	32.60%	31.43%	32.98%	32.93%
Segment	30.00%	27.98%	30.86%	35.41%
Sonar	18.78%	21.08%	22.23%	21.68%
Spiral	44.20%	44.65%	44.52%	44.13%
Statheart	22.72%	23.40%	23.85%	24.22%
Student	4.16%	4.75%	5.24%	5.37%
Wdbc	7.73%	6.95%	6.64%	7.31%
Wine	8.55%	6.59%	9.35%	11.02%
Z_F_S	6.46%	6.66%	6.90%	6.61%
ZO_NF_S	6.01%	6.03%	6.08%	6.25%
ZONF_S	1.79%	1.75%	1.80%	1.79%
ZOO	9.07%	8.63%	7.00%	7.53%
AVERAGE	20.02%	20.12%	20.52%	21.00%

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DATASET a = 1.0a = 1.5a = 2.0a = 4.0**ABALONE** 4.33 4.39 4.46 5.06 AIRFOIL 0.003 0.003 0.002 0.002 BASEBALL 67.45 79.78 74.66 86.54 BK 0.02 0.019 0.019 0.02 BL0.002 0.001 0.001 0.003 CONCRETE 0.003 0.003 0.003 0.003 DEE 0.20 0.20 0.20 0.20 HOUSING 26.24 29.25 26.62 28.13 FRIEDMAN 1.33 1.19 1.20 1.20 FY 0.039 0.04 0.042 0.045 НО 0.014 0.013 0.014 0.014 LASER 0.0027 0.0025 0.0025 0.0024 LW 0.011 0.011 0.013 0.016 MB 0.048 0.049 0.051 0.09 MORTGAGE 0.21 0.31 0.37 0.62 **PLASTIC** 2.20 2.05 2.18 2.21 0.023 0.021 PL 0.023 0.022 PY 0.022 0.023 0.016 0.028 **QUAKE** 0.043 0.038 0.04 0.039 0.024 0.024 0.025 0.026 SN **STOCK** 3.47 3.59 3.68 3.35 TREASURY 0.44 0.42 0.40 1.01 0.029 VE 0.023 0.0240.024**AVERAGE** 4.64 4.92 5.25 5.64

Table 7. Experiments for regression datasets using a variety of values for the margin factor *a*.

Also, the statistical comparisons for these experiments are shown graphically in Figures 5 and 6 respectively.

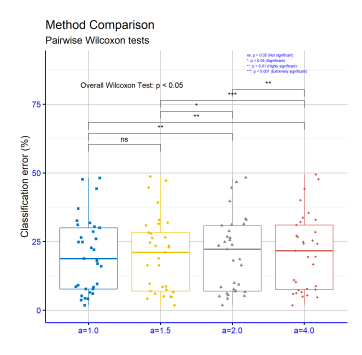


Figure 5. Statistical comparison for the experimental results on the classification datasets, using the proposed method and different values for the margin factor *a*.

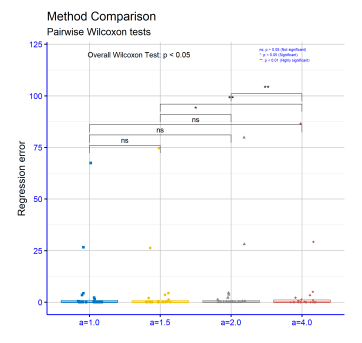


Figure 6. Statistical comparison for the experiments on the regression datasets, using the proposed method and different values for the margin factor *a*.

The parameter a significantly impacts performance, with higher values $(1.0 \rightarrow 4.0)$ correlating with improved mean error rates in both classification $(20.02\% \rightarrow 21.00\%)$ and regression $(4.64 \rightarrow 5.64)$. However, this trend is not uniform. For example, in classification, Segment improves dramatically $(30.00\% \rightarrow 35.41\%, p = 0.0015)$ with increased a, while Lymography degrades $(28.60\% \rightarrow 26.69\%)$. In regression, increasing a introduces instability some problems (e.g., TREASURY: $0.44 \rightarrow 1.01$). Statistical tests confirm significant differences for higher a values (e.g., a = 1.5 vs a = 4.0: p = 0.00065 in classification, p = 0.0021 in regression).

In the evaluation of parameter a for classification datasets (Figure 5), comparisons between higher values of a (e.g., a = 1.5 vs a = 4.0, p = 0.00065) show strong statistical significance, highlighting that increasing a improves performance. Conversely, the comparison a = 1.0 vs a = 1.5 (p = 0.38) reveals no significant difference. For regression datasets (Figure 6), differences become significant mainly at higher a values: a = 1.5 vs a = 4.0 (p = 0.0021) and a = 2.0 vs a = 4.0 (p = 0.0031). However, the comparison a = 1.0 vs a = 4.0 (p = 0.059) approaches the significance threshold (p = 0.05) without crossing it.

4. Conclusions

The study confirms the superiority of the proposed model compared to existing methods in both classification and regression tasks. This superiority is evidenced by a statistically significant reduction in error, though the model's performance is not entirely independent of the data. Instead, it largely depends on data characteristics such as complexity, stratification, or the presence of noise. This underscores the importance of tailoring models to the specificities of each problem, as generalization without careful consideration of specific conditions may lead to degraded results. Additionally, the choice of methods for calculating critical parameters, such as differential weights, appears to have relatively limited impact on overall performance. Differences between approaches (e.g., FIXED, ADAPTIVE, RANDOM) are minimal and often statistically insignificant, indicating that model optimization may require focusing on other factors. At the same time, tuning the parameter *a* emerges as a critical factor: while higher values enhance accuracy in many cases, particularly for large or complex datasets, the same adjustment may introduce instability in applications

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where data sensitivity is high (e.g., financial forecasting). This dual behavior highlights the need to balance flexibility and robustness during the tuning process.

For future research, it would be valuable to explore the role of dynamic algorithms that automatically adjust the parameter a based on data dynamics. For example, machine learning mechanisms that analyze data variability or heterogeneity in real time could optimize the value of a without human intervention, ensuring both stability and high performance. Additionally, the development of hybrid methods that combine the advantages of different approaches (e.g., the adaptability of ADAPTIVE with the simplicity of FIXED) could create more robust solutions capable of addressing a broader range of problems. It is also worth investigating in depth how specific data characteristics—such as noisy measurements, class imbalances, or lack of labeling—affect the model's effectiveness. Such analysis would aid in developing adaptive strategies for data preprocessing or augmentation. Furthermore, the generalizability of findings to more complex environments (e.g., dynamic systems, timeseries data) or different domains (e.g., biomedical data, social networks) remains an open research area. Finally, integrating explainable artificial intelligence (XAI) techniques, such as feature contribution analysis or visualization tools, could enhance model transparency. This would not only facilitate result interpretation by experts but also help identify optimal parameter-tuning practices, transforming the model into a more predictable and reliable tool for real-world applications.

Author Contributions: V.C. and I.G.T. conducted the experiments, employing several datasets and provided the comparative experiments. D.T. and V.C. performed the statistical analysis and prepared the manuscript. All authors have read and agreed to the published version of the manuscript.

Funding: This research received no external funding.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Informed Consent Statement: Not applicable.

Acknowledgments: This research has been financed by the European Union: Next Generation EU through the Program Greece 2.0 National Recovery and Resilience Plan, under the call RESEARCH – CREATE – INNOVATE, project name "iCREW: Intelligent small craft simulator for advanced crew training using Virtual Reality techniques" (project code:TAEDK-06195).

Conflicts of Interest: The authors declare no conflict of interest. Not applicable.

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