Fast Neural Networks Without Multipliers

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Abstract—The paper introduces multilayer perceptrons with weight values restricted to powers-of-two or sum of power-of-two. In a digital implementation, these neural networks do not need multipliers but only shift registers when computing in forward mode, thus saving chip area and computation time. A learning procedure, based on back-propagation, is presented for such neural networks. This learning procedure requires full real arithmetic and therefore must be performed off-line.

Some test cases are presented, concerning MLP's with hidden layers of different size, on pattern recognition problems. Such tests demonstrate the validity and the generalization capability of the method and give some insight into the behavior of the learning algorithm.

I. INTRODUCTION

In the past years, artificial neural networks (NN) have been applied to a variety of problems in many fields, most of them calling for an efficient hardware implementation. Using electronic or optical techniques, several different approaches have been proposed to face this problem. Among them, the digital approach is the most popular. It consists in the numerical simulation of NN's using digital VLSI hardware, parallel computers, or traditional computers. Using this approach, it is possible to take advantage of state-of-the-art VLSI and ULSI implementation techniques, with several advantages over the analog implementation [1].

One of the major problems of digital architectures implementing NN's, affecting both performance and chip area, is the presence of multipliers. In almost every neural model, in fact, the elementary processor (neuron) computes an activation function on the weighted sum of its inputs. While weights are stored and recalled easily in a local memory, the activation function can also be easily computed through and look-up table and sums are performed quickly using a limited amount of hardware; the multiplications between inputs and weights can be the bottle-neck of the system. They are slow compared to other operations and the multipliers require of lot of chip area if a direct VLSI or ULSI implementation is planned. A possible solution to this problem has already been exploited in digital filter design: it consists in avoiding multiplications using integer weights whose values are power-of-two or sums of power-of-two [1], [2]. Using such weights, it is possible to substitute multiplications with simple shifts or shift-and-add operations, which are much faster and require less hardware.

In this paper we demonstrate the feasibility of this approach for the multilayer perceptron (MLP) [3] with binary outputs, and present a learning procedure based on backpropagation (BP) [4] to obtain a MLP with such weights. This learning procedure requires full real arithmetic and therefore must be performed off-line. Once the weights are computed a specific problem, the corresponding power-to-two values can be loaded into the multilayer neural network that will be run in forward mode, processing the data. In the following, after briefly presenting a notation for MLP description, we shall discuss the problem of assigning the neurons to the available processing elements, describe in detail the learning algorithm, and present several test cases concerning MLP's of different sizes and with different numbers of processing elements. These tests confirm the validity of previous results [5], indicate the validity of the method, and give some insight into the behavior and generalization capabilities of the learning algorithm.

II. MULTILAYER PERCEPTRON (MLP)

Many NN models have been proposed in technical literature; among them the MLP is one of the most widely used because a simple and powerful learning algorithm, known as backpropagation has been devised for it. Here we shall briefly describe this model and we shall introduce a notation for its description that will be followed throughout the paper.

In the MLP, all neurons are grouped in sequentially connected layers, numbered from 0 to M. Let $N_k$ be the number of neurons belonging to the generic $k$th layer. Each neuron is connected to all neurons of the two adjacent layers. The neurons of layer 0 (input layer) do not perform computations but only feed input signals into neurons of layer 1. Layer $M$ is the output layer, from where the network response comes.

Each MLP neuron is characterized by one output and many inputs, which are the neuron outputs of the preceding layer. Let $o_{k}(s)$ denote the output of $k$th neuron of the $s$th layer, then the computation performed by each neuron can be expressed as:

$$\text{net}_{k}(s) = \sum_{j=1}^{N_{k-1}} w_{kj}(s) o_{j}(s-1) + \theta_{k}(s)$$  \hspace{1cm} (1)

$$o_{k}(s) = f\left(\text{net}_{k}(s)\right)$$  \hspace{1cm} (2)

where $\text{net}_{k}(s)$ is the weighted sum of the $k$ neuron of the $s$th layer, $w_{kj}(s)$ is the weight by which the same neuron multiplies the output $o_{j}(s-1)$ of the $j$th neuron of the previous...
layer, \( \theta_k(s) \) is the offset of the \( k \)th neuron of the \( s \)th layer and \( f(\cdot) \) is a nonlinear, bounded function, often the sigmoid function.

When processing data, the MLP accepts an input vector \( \vec{i} \) and produces an output vector \( \vec{o} \). From the above considerations, \( \vec{r} \) is equal to \( o^{(0)} \), whose components are \( o_k^{(0)} \), \( k = 1, \ldots, N_0 \). Applying iteratively equations (1) and (2), starting from the first \( (s = 1) \) to the last \( (s = M) \) layer, an output vector \( \vec{o} \) equal to \( o^{(M)} \), is obtained, whose components are \( o_k^{(M)} \), \( k = 1, \ldots, N_M \).

The backpropagation [4] is the most widely used training algorithm in MLP networks. Using an iterative steepest descent technique, it is able to compute the values of the weights and offsets which approximate a desired network behavior. BP requires a number of forward moves of \( o^{(M)} \) for given input vector), each followed by a backward move, during which weight and offset adjustments occur to adapt the network to a desired output vector \( \vec{d} \). This operation can be expressed in its simplest form by the following formulae:

\[
\sigma_k(s) = \begin{cases} 
\frac{d_k - o_k(s)}{\sum_{j=1}^{N_k+1} w_{kj}(s+1) \delta_j(s+1)} & s = M \\
\sigma_k(s) f'(net_k(s)) & s = 1, \ldots, M - 1
\end{cases}
\]  

\[
\delta_k(s) = \sigma_k(s) f'(net_k(s))
\]

\[
\Delta w_{kj}(s) = \eta \delta_k(s) o_j(s-1) 
\]

where \( f'(\cdot) \) is the derivative of the activation function \( f(\cdot) \), \( \eta \) a constant called the learning rate and \( \Delta w_{kj}(s) \) is the weight change. Similar formulae are used to adjust offsets. By applying iteratively equations (3)–(5) from the last \( (s = M) \) to the first \( (s = 1) \) layer, the backward move is completed.

III. NEURONS AND NEURAL NETWORKS WITHOUT MULTIPLIERS

A. Weight Values

The starting point of our approach to the realization of fast NN without multipliers are (1) and (2). The basic activity of a single neuron consists in calculating the weighted sum \( net_k(s) \) of its inputs and then computing the activation function \( f(\cdot) \). The multiplications concern only the weights and the corresponding inputs, not the offset. To avoid numeric multiplications, the weight values must be restricted to power-of-two (or sums of power-of-two).

In this case, the multiplications can be substituted by shift, or shift-and-add operations whose hardware implementation is very simple and much faster than multiplication. The number of bit-shifts that can be performed equals the difference between the maximum and the minimum power-of-two term. In the case of the sum of two power-of-two terms, two shift registers and one adder are needed.

Fig. 1 shows the degree of discretization and nonuniformity of the value distribution for both single power-of-two and sum of two power-of-two terms, with a maximum bit-shift of six.

Throughout the paper, however, we will use a normalization procedure yielding weights, whose absolute values are less than or equal to a given value (usually one).

As they are not involved in multiplications, the offset values can still be real numbers, thus providing more “degrees of freedom” in learning. The dual case in which inputs to neurons are restricted to power-of-two will not be considered in the paper, although it could be a matter for further research.

In the presented results we shall consider the case of a single power-to-two, which represents the most favorable case for hardware implementation, as additional sums are eliminated. It also represents the most difficult case as far as learning is concerned because the discretization is extreme. The learning algorithm proposed in the next section, however, is applicable also to more general NN’s, whose weights are sums of two or more power-of-two terms. Let us denote with \( W_s \) the set of admissible weight values:

\[
W_s \equiv \{ x | x = R \ 2^{-p}; R \in \{-1, 0, 1\}; p \in \{0, 1, \ldots, S\} \}
\]

where \( S \) is the number of bits that can be shifted. It can be noted that this definition constrains the weights absolute value to be less than or equal to one, which is easily obtained by scaling weights and offsets.

Extending admissible weight values to the sum of two power-of-two terms leads to the set:

\[
W_{ST} \equiv \{ x | x = R' 2^{-p} + R'' 2^{-q}; R', R'' \in \{-1, 0, 1\}; \\
p \in \{0, 1, \ldots, S\}; q \in \{0, 1, \ldots, T\} \}
\]

where \( S \) and \( T \) are the maximum number of shifts in each shift register, respectively, and the maximum weights absolute value is less than or equal to two.

It is also worth noting that increasing the number of power-of-two terms of the sum enables the approximation of any number in a given interval to a given error. The expansion in the sum of more than two power-of-two terms has been recently proposed in digital filter design [5], limited to a small subset of weights to fulfill the filter requirements. Such expansion, however, would substantially weaken the advantages of this technique, owing to the increased complexity of weight management and the higher number of operations.
B. Neurons and Processing Elements

In a digital implementation of NN’s, one of the basic problems is the assignment of the neurons to the actual processing elements (PE) that constitute the available digital hardware. Depending on the number of PE’s, the possible choices span from the assignment of a single neuron to each PE to a single PE performing the simulation of the whole NN. At the former extreme there are VLSI implementations of high granularity, while at the latter there are software simulations on a single CPU computer. In the middle there are many possible solutions, implemented directly on VLSI chips, or on many connected microprocessors running in parallel, or on parallel or multi-CPU computers, characterized by assigning the simulation of more than one neuron to every PE with various possible levels of granularity.

In a fast digital implementation the activation function \( f(\cdot) \) is often computed through a look-up table (LUT) contained in every PE. Depending on the granularity of the NN, i.e. on the number of neurons that are assigned to each PE, and on the way neurons are assigned to PE’s, we may consider different cases:

- a distinct LUT for every neuron;
- a LUT common to the neurons of each layer;
- a LUT common to the neurons of each vertical “slice” of the MLP;
- a LUT common to the whole NN.

Fig. 2(a)-(d) graphically show these four possible assignments of MLP neurons to PE’s and thus the possible LUT assignments to neurons. Cases (a) and (d) are straightforward, while cases (b) and (c) are derived by architectures implementing MLP and found in the literature [7], [8]. Of course, other assignments of neurons to PE’s can be found, and the presented learning algorithm can be easily adapted to deal with any possible assignment. Since each LUT allows a distinct scaling of all the weights of the neurons using that LUT, the NN has more degrees of freedom with a greater number of LUT’s. This means a better convergence in learning, as test cases presented in the next sections will show.

IV. LEARNING ALGORITHM

A. Basic Ideas

Given a mapping between input and output spaces of the MLP and a training set of pairs \((i_n, d_n)\) of inputs and outputs representing or sampling the mapping, the problem is to find the right MLP architecture (number of layers and of neurons per layer) and its weights (limited to some \(W_S\) or \(W_{S,T}\) ensemble) and offsets values able to perform the desired mapping with minimum error.

The starting point for the learning algorithm is the solution of the same mapping with a classical MLP having continuous weights, applying BP as the learning algorithm. We shall not address here the problem of finding the right number of layers and neurons per layer, but we shall assume that a continuous solution can be found.

The main ideas behind the proposed learning algorithm are two: the first is to scale the weights of the neurons referring to the same LUT in such a way as to maximize the match between the continuous solution and the discrete one. During this scaling, weight absolute values are also restricted to be less than or equal to one. This approach has been proposed in [9] for digital filter design to find the starting point for discrete minimization of filter coefficients. Since every weight and offset is scaled down by the same constant value, the LUT has to be scaled up accordingly: if weights and offsets are scaled by the common factor \(A\) then the domain of the function to be computed using the LUT has to be scaled by the factor \(A^{-1}\).

The second idea is to perform the learning, starting from the discretized continuous solution obtained by BP, again applying BP. It is well known that the BP algorithm computes the gradient of a quadratic error function that is the sum of the squares of the differences between the actual and desired outputs for each input/output pair of the training set [4]. The BP algorithm used in this phase estimates the gradient of this error function and updates weights and offsets applying a discretized BP formula. Offset values, which are continuous, are freely changed toward the steepest descent direction, while weights are changed only if the learning move prescribes a new value that, rounded to the closer value belonging to \(W_S\) or \(W_{S,T}\), differs from the previous one.

Clearly, a good control of the learning rate \(\eta\) is crucial for the good behavior of the algorithm. In such a way, it is possible to reduce the error function and, hopefully, to reach convergence. We have seen that best results are obtained if the starting point is computed by iterating continuous BP learning up to the highest possible degree of convergence.

B. Formal Description

More formally, given a MLP with a certain number of layers
and neurons per layer and a set (training set) of \( P \) input–output pairs \( \{d_p, a_p; p = 1, \ldots, P\} \), the problem is to find the weight and offset values that minimize the error between desired and computed outputs for all input–output pairs of the training set. Usually, the measures of such error are the average quadratic error \( E_2 \):

\[
E_2 = \frac{1}{PN_M} \sum_{p=1}^{P} |d_p - a_p^{(M)}|^2
\]

and the minmax error \( E_X \):

\[
E_X = \max_{p,k} \left\{ |d_{p,k} - a_{p,k}^{(M)}| \right\}
\]

where \( d_{p,k} \) is the \( k \)th component of \( d_p \) (the desired output of the \( k \)th neuron of the last layer, referring to the \( p \)th input–output pair), and \( a_{p,k}^{(M)} \) is the \( k \)th component of \( a_p^{(M)} \).

In the following we assume that the whole NN is assigned to \( L \) LUT’s and that suffix \( r \) refers to weights and offsets of neurons assigned to \( r \)th LUT, \( r = 1, \ldots, L \). We also assume that the weight values belong to \( W_S \), i.e., they are single power-to-two. The extension to \( W_{S,T} \) (sum of two-power-of-two) is straightforward.

The learning procedure consists in the following step.

1) Make the network learn using the classical BP algorithm until convergence and until no more significative improvement can be achieved. Let \( \theta_r^{(s)} \) be the offset of the \( k \)th neuron of the \( s \)th layer, and \( w_{r,k}^{(s)} \) be the weight related to the link from the \( j \)th neuron of the layer \((s-1)\) to the \( k \)th neuron of the \( s \)th layer of such a converged MLP.

2) Normalize weights: compute the maximum weight, \( W_r \), (in absolute value) of each LUT:

\[
W_r = \max_{r,j,k} \left\{ |w_{r,j,k}^{(s)}| \right\}
\]

where index \( r \) refers to the LUT and indexes \( k \) and \( s \) refer to neurons that use the \( r \)th LUT. Divide all weights and offsets by \( W_r \):

\[
w_{r,j,k}^{(s)} = \frac{w_{r,j,k}^{(s)}}{W_r}, \quad \theta_r^{(s)} = \frac{\theta_r^{(s)}}{W_r}.
\]

In this way, all new weights \( w_{r,j,k}^{(s)} \) will belong to the interval \([-1,1]\).

3) For each LUT of the digital architecture, define the scalar function \( e_r(B) \):

\[
e_r(B) = \max_{r,j,k,s} \left\{ \left| w_{r,j,k}^{(s)} - Q \left( B w_{r,j,k}^{(s)} \right) \right| \right\}
\]

where index \( r \) again refers to the LUT, indexes \( k \) and \( s \) refer to neurons that use the \( r \)th LUT, and the operator \( Q[\cdot] \) denotes rounding to the nearest term belonging to \( W_S \). \( B \) a further scale factor; \( e_r(B) \) is a measure of the match between real weights and their rounded approximations belonging to \( W_S \). Following an approach similar to that proposed in [9] for FIR filters design, \( e_r(B) \) is minimized with respect to \( B \) and new weights and offsets are computed for the value \( B_r \) that minimizes \( e_r(B) \):

\[
w_{r,j,k}^{(s)} = Q \left( B_r w_{r,j,k}^{(s)} \right), \quad \theta_r^{(s)} = B_r \theta_r^{(s)}.
\]

The cumulative scale factor for \( r \)th LUT is thus:

\[
A_r = \frac{B_r}{W_r}
\]

The \( r \)th LUT is then scaled accordingly to yield values corresponding to the nonscaled case.

4) Check the behavior of the neural network having the new weights and offsets \( w_{r,j,k}^{(s)} \) and \( \theta_r^{(s)} \). If the new network performs the desired mapping between inputs and outputs with an error \( E_X \) within a given tolerance, the design procedure ends. If not, the following steps are performed.

5) Starting from the network having weights and offsets \( w_{r,j,k}^{(s)} \) and \( \theta_r^{(s)} \), perform a cumulative BP step [10], adding the contribution of all input–output pairs belonging to the training set and compute for each neuron the quantities \( d_{k}^{(s)} \) and \( D_{kj}^{(s)} \):

\[
d_{k}^{(s)} = \sum_{p=1}^{P} \delta_{p,k}^{(s)} \quad s = M_r, \cdots, 1
\]

\[
D_{kj}^{(s)} = \sum_{p=1}^{P} \delta_{p,k}^{(s)} \phi_{p,j}^{(s-1)} \quad k = 1, \ldots, N_s
\]

\[
 j = 1, \ldots, N_{s-1}
\]

where \( p \) refers to the \( p \)th input–output pair of the training set, \( \phi_{p,j}^{(s-1)} \) is the corresponding output of the \( j \)th neuron of layer \((s-1)\), and \( \delta_{p,k}^{(s)} \) has the meaning defined in (4), referring to the \( p \)th input–output pair of the training set. It is now possible to drop the index \( r \), referring to the LUT, as it does not carry any more information.

6) Change the weights and offsets according to the formulae:

\[
w_{kj}^{(s)} : = Q \left[ w_{kj}^{(s)} + \eta D_{kj}^{(s)} \right] \quad s = M_r, \cdots, 1
\]

\[
\theta_k^{(s)} : = \theta_k^{(s)} + \eta d_{k}^{(s)} \quad k = 1, \ldots, N_s
\]

where the right-hand-side term refers to the old value, while the left one is the new value. If the error \( E_2 \) is decreased after application of (16), \( \eta \) is increased by a factor \( M_\eta \). On the contrary, if \( E_2 \) is increased, \( \eta \) is decreased by a factor \( D_\eta \). In this way, \( \eta \) is automatically adapted to the problem, similarly to the approach proposed in [10]. The values of \( \eta \) control parameters actually used are:

\[
M_\eta = 1.05, \quad D_\eta = 0.7.
\]

This learning rate control strategy (a slight increase of \( \eta \) in case of success and a bigger decrease in case of failure) is fairly robust, and has been successfully tested by the authors, also in continuous BP learning. As the learning rate is self-adaptive, the actual values of \( M_\eta \) and \( D_\eta \) are not critical, provided that they do not differ too much from the given ones.

Fig. 3(a) shows a typical learning curve of the method \( E_2 \) versus the number of iterations), while Fig. 3(b) shows the
having the highest values of $|D_{bkj}^{(s)}|$, according to (15) and (16).

b) Change the network topology, adding layers and/or neurons to existing layers, trying to find a better starting point for the discretization procedure.

However, convergence cannot be guaranteed, as in MLP’s with continuous weights using the BP algorithm. The following section shows indeed that convergence has been achieved in most test cases considered.

V. RESULTS

The design procedure has been tested using a pattern recognition problem, consisting in the recognition of character patterns encoded in a $8 \times 8$ pixel matrix, according to the IBM PC CGA character set. 95 different characters had to be recognized, corresponding to ASCII codes between 32 and 126. The number of input–output pairs, $P$, is thus 95, and for each pair the input is a vector of 64 binary values, corresponding to the pixel matrix representing a character, while the output is a vector of 8 binary values, representing its coded ASCII value. Therefore, the used MLP’s have 64 inputs and 8 outputs.

All performed tests deal with binary values, while BP requires real values. We set all input–output pairs of values of the training set to 0.1 and 0.9 rather than to zero and one, respectively, to improve convergence [11].

Let us note that, although the CGA character set has been designed to maximize the difference between characters, there are still many characters very similar to each other. For instance: colon and semicolon; full stop and comma, the characters 1, I, [, :, and many others. Therefore, this pattern recognition problem is not straightforward.

A reduced test problem was also used, consisting in the recognition of the characters representing the ten digits. In this case, only four network outputs are significant.

In the tests, we considered various MLP’s, with various topologies, various assignments of LUT’s to neurons, and different values of $S$ (see (6)).

A. MLP’s Used in the Tests

In the tests we used two MLP’s: the first with 64 inputs, 64 hidden neurons and 8 outputs (64/64/8) and the second with 32 hidden neurons (64/32/8). The assignments of PE’s (and thus LUT’s) to neurons, as discussed in Section III-B, are of four kinds: single ($S$), slice ($C$), layer ($L$), and global ($G$). In the case ($C$), we have 8 slices, each including 8 inputs, 8 or 4 hidden neurons and one output neuron. The considered values of the number of shifts, $S$, are 8, 4, and 1.

For the reduced problem with 10 input–output pairs, we used three different MLP topologies (with 64/64/4, 64/8/4 and 64/4 neurons, respectively), and many different values of $S$, from 8 to 0.

B. Continuous Learning

Step 1 of the learning algorithm prescribes to perform a continuous BP learning, until no further significant improve-
### Table I

<table>
<thead>
<tr>
<th>MLP name</th>
<th>Neurons in Layers</th>
<th>Input</th>
<th>Hidden</th>
<th>Output</th>
<th>approx. n</th>
<th>pairs of iterations</th>
<th>$E_2$</th>
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<tbody>
<tr>
<td>64/64/8 A</td>
<td>64</td>
<td>64</td>
<td>8</td>
<td>95</td>
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<td>0.0299</td>
<td></td>
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<tr>
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<td>64</td>
<td>8</td>
<td>95</td>
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<tr>
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<td>64</td>
<td>64</td>
<td>8</td>
<td>95</td>
<td>10000</td>
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<td></td>
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<tr>
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<td>4</td>
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<td>400</td>
<td>0.0325</td>
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### Table II

<table>
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<th>LUT Assignment to Neurons</th>
<th>Value of $S$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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</tr>
<tr>
<td>64/64/8</td>
<td>avg.</td>
</tr>
<tr>
<td>Single (S)</td>
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<tr>
<td>Slice (C)</td>
<td>107</td>
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<tr>
<td>Layer (L)</td>
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<tr>
<td>Global (G)</td>
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<tr>
<td>64/32/8</td>
<td>avg.</td>
</tr>
<tr>
<td>Single (S)</td>
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<tr>
<td>Slice (C)</td>
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<tr>
<td>Layer (L)</td>
<td>216</td>
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<tr>
<td>Global (G)</td>
<td>194</td>
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</table>

### Table III

<table>
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<th>N. of iterations for $E_X = 0.3$</th>
<th>Starting value of $E_X$</th>
</tr>
</thead>
<tbody>
<tr>
<td>LUT: S</td>
<td>C</td>
</tr>
<tr>
<td>64/64/8; $S = 4$</td>
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<tr>
<td>A</td>
<td>206</td>
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<tr>
<td>B</td>
<td>82</td>
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<td>C</td>
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<td>D</td>
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<td>212</td>
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<tr>
<td>F</td>
<td>128</td>
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<td>G</td>
<td>201</td>
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<td>64/32/8; $S = 8$</td>
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<td>F</td>
<td>382</td>
</tr>
<tr>
<td>G</td>
<td>180</td>
</tr>
</tbody>
</table>

### C. Results with 95 Characters

In this tests, the tolerance $\epsilon$, as defined in point 4) of Section IV-B, was set to 0.3, meaning that a value less than 0.4 is considered to be zero, and a value greater than 0.6 is considered to be one. Table II summarizes all test runs, reporting the average number of iterations and their standard deviation versus $S$, for all LUT assignments to neurons. Table III reports the starting value of $E_X$ and the total number of iterations to converge for all test runs concerning the case with $S = 4$.

The results concerning the 64/64/8 MLP are very noisy, being characterized by a high standard deviation. However, convergence was always achieved in a number of iterations ranging from 48 to 472, lowering the absolute error $E_X$ from an initial value of about 0.8 to less than 0.3. All kinds of LUT assignments to neurons seem to be equivalent, within one standard deviation, in the number of iterations, at least for $S$ greater than one. Moreover, the number of iterations needed to converge seems lower for $S = 4$ than for $S = 8$, while one could expect the contrary, as the number of degrees of freedom decreases with $S$. Therefore, these results confirm the redundancy of the 64/64/8 MLP for this problem.

On the contrary, the results concerning the 64/32/8 MLP, limited to three tests for each value of $S$, show a clear dependence of the MLP behavior on its degrees of freedom given by LUT assignment and shift levels. Convergence in the case with $S = 8$ is much better than in the case with $S = 4$, while with $S = 1$, the MLP's always fail to converge.

These results suggest that, as redundancy becomes critical, convergence becomes slower and cannot be reached with extreme discretization.

These results, however, confirm the validity of the proposed algorithm and the realizability of NN's with power-of-two weights. In the case of extreme weight discretization, of course, a redundant MLP is needed to achieve convergence.

### D. Asymptotic Learning

Beside the ability of the algorithm to reach prescribed value of $E_X$, it is also interesting to find the smallest reached value of $E_X$. As these tests need a lot of computer time, we performed most of them using the reduced test cases with 10 input/output pairs. The used MLP's span from a very redundant one (64/64/4 MLP) to the least redundant possible (64/4/4 MLP). In the tests, we considered also the extreme
discretization with $S = 0$, meaning that weights values must belong to the set $\{-1, 0, 1\}$.

Fig. 4(a)-(c) show the results for the MLP’s 64/64/4, 64/8/4, and 64/4, respectively. Each figure reports the value of the minimum reached $E_X$ versus $S$, for the various assignments of LUT’s to neurons. It can be seen that, when the redundancy of the MLP is high (high number of neurons and/or high value of $S$), the minimum value of $E_X$ reached is approximately the same. As the redundancy diminished, the minimum value of $E_X$ increases and becomes higher as $S$ and as the number of LUT’s decreases, with the exception of the “layer” assignment ($L$), that performs in most cases better than the “slice” one ($C$), which has more LUT’s. Presently, we have no explanation for this fact.

Table IV shows the results of similar test cases, for the 64/64/8 MLP with 95 characters, which confirm this view.

<table>
<thead>
<tr>
<th>LUT Assignment to neurons</th>
<th>$E_X$ Value of $N$</th>
<th>$E_Y$ Value of $N$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single (S)</td>
<td>0.207</td>
<td>0.226</td>
</tr>
<tr>
<td>Single (S)</td>
<td>0.056</td>
<td>0.060</td>
</tr>
<tr>
<td>Global (G)</td>
<td>0.280</td>
<td>0.252</td>
</tr>
<tr>
<td>Global (G)</td>
<td>0.056</td>
<td>0.067</td>
</tr>
</tbody>
</table>

VI. GENERALIZATION CAPABILITY

To perform further tests on the performance of the proposed algorithm, and to verify the generalization capability of the discretized MLP, another experiment has been carried out. This experiment concerns the well known problem of classifying points inside and outside a circular region in the plane, as reported in [12].

Formally, the problem consists in the building of a MLP with 2 inputs and 1 output, which, given the $x, y$ coordinates of a point in the Cartesian plane, is able to recognize if this point lies inside or outside a unit-radius circle centered at the origin. The network output is zero if the point lies outside, one if it lies inside. In practice, the experiment is performed choosing a suitable MLP topology, creating a finite-size learning set composed of randomly selected points inside and outside the circle, and applying the BP algorithm until the network converges within a given accuracy. As in [12], a MLP with one hidden layer of 8 neurons has been used in this experiment, while the learning set is composed of 200 points, 100 points randomly selected inside the circle, and 100 points randomly selected outside the circle, but within the square region bounded by $|x| < 5$ and $|y| < 5$.

To test the generalization capabilities of the resulting network, the MLP has been requested to classify all the points inside the region described by $|x| < 5$ and $|y| < 5$. In practice, a dense grid of points was used to this purpose. The obtained set of output values can be used to produce a contour plot, where each curve corresponds to a given output level. Any point inside the curve produces an output level which is equal to, or greater than the given level. A typical plot for the MLP after convergence is reported in Fig. 5, where the four curves represent the levels 0.2, 0.4, 0.6, and 0.8 from the outer to the inner. As can be seen, the contour plots resemble very much the target unit circle, especially the 0.8 level innermost curve.

The proposed algorithm has been tested on this problem varying LUT assignment and discretization levels. For each
LUT assignment and discretization level, many trials were performed with different initial sets of weights. In all test runs, a good accordance between the overall degree of freedom (LUT number and discretization levels) and the generalization capability of the network has been obtained.

For the sake of brevity, only a few graphical results are reported. Fig. 6(a) shows the performance of the MLP discretized using a global LUT (the worst case of LUT assignment) and one power-of-two term with $S = 0$ (the worst case of level discretization, the weights being constrained to be $-1$, $0$, or $1$). Although the network behavior is worse than that of the original MLP (see Fig. 5), the discretized network obtained with the proposed algorithm retains a good generalization capability as far as the inner levels are concerned. A comparison of this plot with Fig. 6(b) which reports the performance of the network after steps 2) and 3) of the algorithm, but before the additional discretized BP procedure of steps 5) to 7) clearly highlights the importance of this phase to the overall performance.

Fig. 7 reports the result of the MLP discretized using a global LUT and a single power-of-two term with $S = 3$. The generalization capability and the effectiveness of the proposed algorithm are confirmed.

VII. DISCUSSION AND CONCLUSIONS

In this paper, we proposed the use of weights with values limited to power-of-two for a fast digital implementation of NN’s, and presented a learning algorithm for MLP’s with such weights. The proposed discretized MLP and the learning algorithm performed well in all test cases considered and demonstrated a good generalization capability. The main drawback of the algorithm is the need for off-line computation, since learning requires multiplications.

A discrete power-of-two MLP is, of course, less powerful than a classical, real-valued MLP of the same size. The higher the number of admissible power-of-two values, and the more different LUT’s are available, the more powerful the discrete MLP will be. These considerations are confirmed by the tests made: if the real-valued MLP is not redundant at all for solving a given problem, its power-of-two counterpart will exhibit a difficult convergence, if not at all. If the real-valued MLP is redundant, the power-of-two counterpart exhibits a
clear improvement in its performance, increasing the number of admissible power-of-two, \( S \), and the number of LUT's. Eventually, when the real-valued MLP is very redundant, the corresponding power-of-two network shows the typical learning behavior of very redundant MLP's: the number of iterations needed to learn is very variable and often worse than for a less redundant network. In this latter case, discrete MLP's exhibit a limiting behavior that does not depend on \( S \) or on the LUT number, being affected by "noise" due to discretization. On the other hand, a MLP which is redundant enough can learn a given behavior using only one or a few LUT's, even with extreme weight discretization (\( S = 0 \)).

In the presented tests we concentrated on single power-of-two weights, although the learning algorithm can deal also with multiple power-of-two. In fact, we also performed some tests on the character recognition problem using MLP's with weights whose values were the sum of two power-of-two terms, and \( S \) was varied between 7 and 9. Such weight values are much closer to continuous ones than single power-of-two weights, and we could expect a discretized MLP behavior very similar to a real-valued MLP. All test cases computed confirmed this view, presenting a much better convergence to the solution, and often showing a satisfactory behavior immediately after weight normalization and rounding, with no need to perform further learning iterations.

The idea of using power-of-two weights to avoid multiplications in NN's was exploited also for the Neocognition neural model [1]. Owing to Neocognition characteristics, it was possible to avoid multiplications also in the learning phase, thus taking full advantage of this technique.

Reference [13] reports a analysis of the effects of quantization on learning and generalization in NN's. The test cases presented there refer to a classification problem, where the performance of the network is measured in terms of the percentage of patterns correctly classified. However, the qualitative relationship among quantization weight level, number of hidden units (network redundancy), and network performance is similar to that previously presented and discussed.

Other approaches to NN's with discrete weights are reported in [14], where an analog VLSI model of MLP (feedforward and recurrent) with discrete weights is presented, together with a discrete learning algorithm (weight perturbation); and in [15], where extremely discretized NN's with binary weights are proposed, together with a learning procedure (CHIR) based on choosing internal representations. These approaches are not easily comparable with our approach, since in essence they are neural models not based on back propagation. Moreover, the former is based on an analog VLSI implementation, while our proposal is aimed at a digital implementation for the reasons explained in the introduction.

In conclusion, we believe that the power-of-two weight MLP could be good VLSI implementation alternative for applications requiring extremely fast data processing, but not requiring in-line learning or relearning. Although restricted to feed-forward processing, the area-delay product of such neural VLSI chips could be improved by two orders of magnitude, or more, over traditional implementations using multipliers [1]. Examples of possible fields of application could be the same as power-of-two digital filters: DSP of wide-band signals, nonlinear channel equalization, real-time image processing, and others.

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REFERENCES


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