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# Benefits of Gain: Speeded Learning and Minimal Hidden Layers in Back-Propagation Networks

John K. Kruschke and Javier R. Movellan

Abstract - The gain of a node in a connectionist network is a multiplicative constant that amplifies or attenuates the net input to the node. The objective of the work is to explore the benefits of adaptive gains in back propagation networks. First it is shown that gradient descent with respect to gain greatly increases learning speed by amplifying those directions in weight space that are successfully chosen by gradient descent on weights. Adaptive gains also allow normalization of weight vectors without loss of computational capacity, and we suggest a simple modification of the learning rule that automatically achieves weight normalization. Finally, a method for creating small hidden layers by making hidden node gains compete according to similarities between nodes, with the goal of improved generalization performance, is described. Simulations show that this competition method is more effective than the special case of gain decay.

# I. THE GAIN PARAMETER IN BACK PROPAGATION

The back propagation learning algorithm [1]-[5] has become a very popular method for training connectionist networks. Two of the appealing properties of back propagation are its tolerable learning speed and its ability to generalize to novel inputs. Unfortunately back propagation is sometimes too slow, and

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generalization is not always good. In this article we introduce a new parameter, gain, into back propagation networks and show that it can yield benefits for learning speed and generalization.

Consider a multilayer feed-forward network, as in standard back propagation. Let  $a_i^s$  be the activation of the *i*th node of layer s, and let  $a^s = [a_1^s \cdots a_n^s]^T$  be the column vector of activation values in layer s. The input layer is layer 0. Let  $w_{ij}^s$  be the weight on the connection from the *j*th node in layer s-1 to the *i*th node in layer s, and let  $w_i^s = [w_{i1}^s \cdots w_{in}^s]^T$  be the column vector of weights from layer s - 1 to the *i*th node of layer s. The net input to the *i*th node of layer s is defined as  $net_i^s =$  $\langle \boldsymbol{w}_{i}^{s}, \boldsymbol{a}^{s-1} \rangle = \sum_{k} w_{ik}^{s} a_{k}^{s-1}$ , and let  $\mathbf{net}^{s} = [\mathbf{net}_{1}^{s} \cdots \mathbf{net}_{n}^{s}]^{T}$  be the column vector of net input values in layer s. The activation of a node is given by a function of its net input,

$$a_i^s = f(g_i^s \operatorname{net}_i^s), \tag{1}$$

where f is any function with a bounded derivative, and  $g_i^s$  is a real number called the gain of the node.

Suppose that for a particular input pattern,  $a^0$ , the desired output is the teacher pattern  $t = [t_1 \cdots t_n]^T$ , and the actual output is  $a^{L}$ , where L denotes the output layer. Define an error function on that pattern,  $E = (1/2)\sum_i (t_i - a_i^L)^2$ . The overall error on the training set is simply the sum, across patterns, of the pattern error E. We then perform gradient descent on Ewith respect to  $w_{ij}^s$ . The chain rule yields

$$\frac{\partial E}{\partial w_{ij}^s} = \frac{\partial E}{\partial \mathbf{net}^{s+1}} \cdot \frac{\partial \mathbf{net}^{s+1}}{\partial a_i^s} \cdot \frac{\partial a_i^s}{\partial \mathbf{net}_i^s} \cdot \frac{\partial \mathbf{net}_i^s}{\partial w_{ij}^s}$$
$$= \left[ -\delta_1^{s+1} \cdots - \delta_n^{s+1} \right] \cdot \begin{bmatrix} w_{1i}^{s+1} \\ \vdots \\ w_{ni}^{s+1} \end{bmatrix} \cdot f'(g_i^s \operatorname{net}_i^s) g_i^s \cdot a_j^{s-1}, \quad (2)$$

where  $\delta_i^s = -\partial E / \partial \operatorname{net}_i^s$ . In particular, the first three factors of (2) indicate that

$$\delta_i^s = \left(\sum_k \delta_k^{s+1} w_{ki}^{s+1}\right) f'(g_i^s \operatorname{net}_i^s) g_i^s.$$
(3)

The recursive formula (3) for  $\delta^s$  is the same as in standard back propagation [1], [2] except for the appearance of the gain parameter. Combining (2) and (3) yields the learning rule for weights:

$$\Delta w_{ii}^s = \epsilon_w \delta_i^s a_i^{s-1},\tag{4}$$

where  $\epsilon_w$  is a small positive constant called the "step size" of gradient descent with respect to weights.

Gradient descent on error with respect to the gains can also be computed. Using the chain rule as previously, it is easy to compute that

$$\frac{\partial E}{\partial g_i^s} = \left(\sum_k \delta_k^{s+1} w_{ki}^{s+1}\right) f'(g_i^s \operatorname{net}_i^s) \operatorname{net}_i^s.$$
(5)

Then

$$\Delta g_i^s = \epsilon_g \delta_i^s \operatorname{net}_i^s / g_i^s, \tag{6}$$

where  $\epsilon_g$  is the step size of the gains. The learning rule for gains (6) is easily incorporated into standard back propagation programs. In particular, all the quantities that appear in (6) are locally available to the affected gain  $g_i^s$ .

An equivalent method was first introduced by Movellan [6] and independently proposed by Tawel [7]. Other authors (e.g.,

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RATIOS OF A	Average L	EARNING	TIMES (IN SWEEP	s) for BP Rela	TIVE TO BPG <sup>a</sup>
Number			Criterion	= 0.20	
Hidden			Weight Step	Size, $\epsilon_w$	
Nodes	0.25	0.5	1.0	2.0	4.0
2	1.9	1.7	2.1(1/1)	1.3(1/2)	2.3(8/11)
4	1.3	1.4	1.8	1.8	5.3
8	1.5	1.4	1.5	1.3	1.4
16	1.5	1.4	1.3	1.3	1.2
32	1.6	1.4	1.3	1.3	1.2
Number			Criterion	= 0.10	
Hidden			Weight Step	Size, $\epsilon_w$	
Nodes	0.25	0.5	1.0	2.0	4.0
2	3.0	2.4	2.4(1/1)	1.4(1/3)	3.5(8/11)
4	1.9	1.8	2.0	1.9	4.9
8	1.9	1.5	1.7	1.4	1.3
16	2.0	1.6	1.4	1.4	1.2
32	2.2	1.7	1.5	1.3	1.2
Number			Criterion	= 0.05	
Hidden			Weight Step	Size, $\epsilon_{}$	
Nodes	0.25	0.5	1.0	2.0	4.0
2	5.7	5.1	4.3(1/1)	1.6(1/3)	3.5(8/11)
4	4.2	3.7	2.9	2.2	4.7
8	4.1	2.6	2.2	1.5	1.4
16	4.2	2.8	1.7	1.5	1.3
32	4.6	3.3	1.8	1.4	1.3

TABLE I

<sup>a</sup>When applied to the 4-4 encoder problem. Numbers in parentheses indicate the number of times (out of 15) that BP or BPG failed to reach criterion in 1000 sweeps.

[8]-[10]) have also used a gain parameter, with different purposes and different architectures. These are discussed later.

Throughout this article, we will refer to back propagation with adaptive gains as BPG, and to standard back propagation, for which  $g_i^s = 1$  for all *i*, *s* and  $\epsilon_g = 0.0$ , as BP.

### II. Speeded Learning from Gradient Descent with Gain

Learning time in BPG is remarkably faster than standard BP. In this section we discuss some criteria for comparing learning times of different algorithms, and then report our simulation procedure and results.

### Comparing Learning Speeds

Comparing the speeds of variants of back propagation can be tricky. One must control all of the following variables: updating procedure (every trial vs. periodic), solution criteria, initialization procedure, learning rates, number of hidden nodes, and error function. In our comparisons of BP and BPG, we performed gradient descent on the sum of squared errors, weight and gain changes were accumulated after each pattern presentation but only executed at the end of a full sweep (or "epoch") through the training set, and learning time was measured as the number of sweeps until  $|t_i - a_i^L| < ecrit$ , for a pre-set criterial value *ecrit*, across all patterns and output nodes *i*.

Since the initial values of the weights may affect convergence, it is common practice to run several simulations with different starting weights and to combine the results with appropriate statistics. The choice of these statistics is not straightforward. Arithmetic averages are excessively influenced by the occasions when the network fails to reach criterion in the allotted time, but robust statistics, like the median, do not reflect those occasions at all. In our simulations we characterized speed with two statistics: the arithmetic average of the number of sweeps when the solution was achieved in less than 1000 sweeps; and, the number of simulations when the system needed more than 1000 sweeps.

Combining the results for different step sizes is also tricky. One possibility is to compare learning times for optimal stepsizes. However, it is often preferable to learn moderately fast over a wide range of step sizes than to learn exceptionally fast over a very limited range. In this article we report learning times for a variety of parameter combinations. We also report averages of the different combinations to give an estimate of the relative efficiencies of the algorithms.

### Procedure

We compared BP and BPG on two standard benchmark problems, the exclusive-or (XOR) and the 4-4 encoder problem [1]. We tested learning times of BP and BPG on all combinations of the following independent variables.

- Number of hidden nodes: 2, 4, 8, 16, 32.
- Step size  $\epsilon_w$  for weight change: 0.25, 0.5, 1, 2, 4.
- Criterial value (ecrit) for solution: 0.20, 0.10, 0.05.

There was a single hidden layer, in which all nodes used the logistic activation function  $a = 1/(1 + \exp(-g \operatorname{net}))$ . In each condition 15 different initializations for weights were used, with weights assigned according to a uniform random distribution from -1 to +1, such that the fan-in weight vector (including bias) of each node was normalized to Euclidean length 1. The step sizes for weights in the output layer were scaled according to the the standard fan-in correction formula (Plaut *et al.*, [10]).

In BPG, gains were modified according to (6) with  $\epsilon_g = 0.20$ . In the XOR problem, gain was restricted such that 0.75 < g < 3.50, so that gains would be neither too small, making the node ineffective, nor too large, producing excessive change. Momentum (see [1]) was set to 0.9 for both weights and gains. To assure that our program was correct, its performance, with  $\epsilon_g = 0.0$  and g = 1, was compared to McClelland and Rumelhart's [11] backpropagation software. The programs behaved identically to the fifth decimal place.

T T

RATIOS	OF AVERAGE L	EARNING TIMES	(IN SWEEPS) FOR	BP RELATIVE T	o BPG <sup>a</sup>
Number Hidden		W	Criterion = 0.20 eight Step Size,	€w	
Nodes	0.25	0.5	1.0	2.0	4.0
2	1.2(7/3)	1.4(0/2)	3.2(0/1)	1.8(0/2)	1.2
4	1.2(2/2)	2.0	3.8	2.7	1.4
8	1.6(2/2)	1.5	2.3	3.1	1.6
16	1.4(4/1)	1.3(1/0)	1.8	3.0	3.0
32	1.3(8/2)	1.4	1.4	1.9	3.6
Number			Criterion $= 0.10$		
Hidden		W	eight Step Size,	e,,,	
Nodes	0.25	0.5	1.0	2.0	4.0
2	1.3(7/3)	1.5(0/2)	3.3(0/1)	1.8(0/2)	1.3(0/1)
4	1.3(4/2)	2.1	3.9	2.8	1.4
8	1.7(3/2)	1.6	2.5	3.1	1.6
16	1.5(6/1)	1.5(1/0)	2.0	3.1	3.1
32	1.2(11/3)	1.6(2/0)	1.6	2.2	3.7
Number			Criterion $= 0.05$		
Hidden		W	eight Step Size,	Ew	
Nodes	0.25	0.5	1.0	2.0	4.0
2	1.5(8/3)	1.9(0/2)	3.5(0/1)	1.8(0/2)	1.3(0/1)
4	1.5(8/2)	2.5(1/0)	4.5	3.0	1.5
8	1.8(10/2)	2.0(2/0)	3.2	3.5	1.7
16	1.4(4/3)	1.8(4/0)	2.6	3.8	3.4
32	1.3(15/7)	1.6(9/0)	2.3	3.0	4.2

TABLE II

<sup>a</sup>When applied to the XOR problem. Numbers in parentheses indicate the number of times, out of 15, that BP/BPG failed to reach criterion within 1000 sweeps.

RATIOS OF	TABLE III Ratios of Learning Times for BP and BPG <sup>a</sup>			
Criterion	Encoder	XOR	Average	
0.05	2.9	2.4	2.7	
0.1	1.9	2.1	2.0	
0.2	1.7	2.0	18	

2.2

2.2

<sup>a</sup>Summarized from Tables II and III.

Average

2.2

#### Results

Tables I and II show the learning times of BP relative to BPG for the 4-4 encoder problem and the XOR problem. Average results are summarized in Table III. For all trials when criterial error was reached, BPG was faster than BP. The learning time ratios vary from 1.1 to 5.7, with an average of 2.2. In terms of the number of times that criterial error was not reached in 1000 sweeps, BPG also performed better than BP except when two hidden nodes were used. In that case BPG could not solve the problem 13% of the time, as opposed to BP that did not solve it 11% of the learning criteria and the type of problem. The average learning time ratio is 1.83 for a criterion of 0.2, and 2.68 for a criterion of 0.05. In a previous study [6] it was shown that for smaller criteria, BPG is more than 6 times faster than BP.

# III. DISCUSSION

### Speeded Learning by Other Methods

There have been many other back propagation speed-up methods described in the literature, e.g., Fahlman's [12] Quick-Prop, or Jacob's [13] learning-rate adjustment heuristics. We do not claim that BPG is faster or incompatible with these methods. An advantage of adaptive gain is that it is easy to introduce into back propagation programs, and that it accelerates learning without the need to invoke principles other than gradient de scent. Adaptive gains also provide a variety of other benefits that we will discuss in later sections.

### Relation of Gain Change and Weight Change

In this section we explore the relationship between gain and weight change. We then analyze why BPG speeds up learning relative to BP.

Lemma 1: Suppose weights are changed by gradient descent on the error E, and gains are also changed by gradient descent on E. Then, for a given node, the magnitude of its gain increases if and only if the length of its weight vector increases.

*Proof:* Gradient descent on the error E with respect to weights and gains can be expressed as follows (cf. (2)–(6):

$$\Delta \boldsymbol{w}_{i}^{s} = \boldsymbol{\epsilon}_{w} \frac{\partial E}{\partial (g_{i}^{s} \operatorname{net}_{i}^{s})} g_{i}^{s} \boldsymbol{a}^{s-1}$$
(7)

$$\Delta g_i^s = \epsilon_g \frac{\partial E}{\partial (g_i^s \operatorname{net}_i^s)} \langle w_i^s, a^{s-1} \rangle \tag{8}$$

Solving (7) for  $a^{s-1}$  and substituting into (8) yields

$$\Delta g_i^s = \frac{\epsilon_g}{\epsilon_w} \frac{1}{g_i^s} ||\mathbf{w}_i^s|| \, ||\Delta \mathbf{w}_i^s|| \cos\theta \tag{9}$$

where  $\theta$  is the angle between  $w_i^s$  and  $\Delta w_i^s$ . Note also that

$$\Delta \|\mathbf{w}_{i}^{s}\|^{2} = \|\Delta \mathbf{w}_{i}^{s}\|^{2} + 2\|\mathbf{w}_{i}^{s}\|\|\Delta \mathbf{w}_{i}^{s}\|\cos\theta.$$
(10)

Case I:  $\cos \theta \ge 0$ . From (9),  $\Delta g_i^s \cdot g_i^s > 0$ . Therefore, the signs of  $g_i^s$  and  $\Delta g_i^s$  are the same, so  $\Delta |g_i^s| > 0$ . From (10),  $\Delta ||\mathbf{w}_i^s||^2 > 0$ . In this case, then, the signs of  $\Delta ||g_i^s|$  and  $\Delta ||\mathbf{w}_i^s||$  agree: They both are positive.

Case II:  $\cos \theta < 0$ . From (9),  $\Delta |g_i^s| < 0$ , by argument analogous to that used in Case I. From Equation 10, we can see that  $\Delta ||w_i^s||^2 < ||\Delta w_i^s||^2$ . For arbitrarily small<sup>1</sup> step sizes,  $||\Delta w_i^s||^2 \to 0$ , so that  $\Delta ||w_i^s||^2 < 0$ . Again the signs of  $\Delta |g_i^s|$  and  $\Delta ||w_i^s||$  agree: They both are negative.

Adaptive gain has a catalytic effect in the learning process by modifying the magnitude, not the direction, of the weight change. Empirical observations show that in the early phases of learning, when a successful direction has not yet been found, weight vectors change direction from trial to trial and both gain and weight-vector length shrink. As successful directions are found, weights and gains grow larger. From (3) and (4), gain can be seen as modulating the stepsize  $\epsilon_w$  of the weight change, amplifying learning in nodes that find successful directions in weight space. This may help to explain why BPG does not necessarily provide a big advantage with very small networks: Larger networks provide a larger pool of candidate directions, so that the benefits of gain are more likely to be utilized.

# Gain and Weight Normalization

By including the gain parameter in back propagation, one can also normalize weight vectors without loss of computational power. One advantage of normalization is that it maintains the weight values within bounds [14], [15]. Normalization can also be used to facilitate comparison between nodes [16]. Weight vectors can be factored into length and direction, with direction indicated by the normalized weight vector, and length indicated by the gain of the node. By performing gradient descent on gain, and by implication from Lemma 1, the effects of back propagation on the lengths of the weight vectors will be reflected by the gains, even with the weight vectors themselves normalized.

Several approaches might be used to normalize the weight vectors. One possibility is to let the weights change via back propagation and then to normalize the weights directly, dividing them by the length of the weight vector. Such an approach was taken by Kruschke [16]. Another possibility, used by Oja [15] in the context of a Hebbian rule, is to generate a Taylor series approximation to the normalization process. Movellan [17] derived a Taylor series approximation to normalization for back propagation:

$$\Delta w_{ij}^s = \epsilon_w \delta_i^s \left( a_j^{s-1} - \operatorname{net}_i^s w_{ij}^s \right), \tag{11}$$

where  $\Delta w_{ij}^s$  is the modified weight change. Notice that the normalized rule of (11) is nearly the same as the standard weight update rule of (4) but with a corrected activation from the layer below. An advantage of the Taylor approximation over "brute-force" normalization is that the length of the weight vector need not be computed. We have applied the normalization rule of (11) in conjunction with BPG, with good results [17].

#### IV. MINIMAL HIDDEN LAYERS FROM GAIN COMPETITION

We have shown that the gain parameter is useful for speeding up learning and for allowing weight vector normalization. We will now show that it can also be used to create hidden layer bottlenecks; i.e., hidden layers with relatively few participating nodes. Small hidden layers have been shown to improve the generalization performance of back-propagation networks in some applications.

# Gain Indicates Participation in Representation and Learning

We say that a given hidden node *participates in representing* the input if and only if its activation changes for some change of input. The gain of a node indicates how much the node participates in representing the input. When the gain of a node is zero, its activation is constant: If  $g_i = 0$ , then  $a_i = f(g_i \text{ net}_i) = f(0)$  for all net<sub>i</sub>, and as the magnitude of the gain grows, the variance of activation values from f(0) increases for monotonic f.

The gain of a node also indicates the extent to which the node participates in learning. When the gain of a node is zero, the error propagated to its weights is also zero, and hence its weights cannot learn. When the magnitude of the gain grows, the learning by weights also increases. That can be seen explicitly from (3), in which gain acts as a continuous modulator of the magnitude of error propagated to the weights.

The gain parameter is not just a passive indicator like the pointer of a speedometer. If one tries to go faster by moving the pointer of a speedometer to a higher value, one only succeeds in breaking the speedometer. On the contrary, if we adjust the gain of a node, we actually change its activation variance and its weight learning rate. It is the causal efficacy of the gain parameter that makes it an attractive device for introducing constraints into back propagation.

### Improved Generalization from Small Hidden Layers

There is empirical evidence that generalization to novel input patterns is improved by using hidden layers with a small number of nodes (e.g. [18]-[20]). In these cases, generalization from the training set to novel inputs was better when the number of hidden nodes was relatively small.<sup>2</sup> The reason for improved generalization is intuitively clear: A small hidden layer forces the input patterns to be mapped through a low-dimensional space, enforcing proximities between hidden-layer representations that were not necessarily present in the input-pattern representations. Only the differences between patterns that are most important for decreasing error will be preserved as large distances between hidden layer patterns. Differences between input patterns that are not preserved in the hidden layer representation are thereby generalized over completely. A more detailed discussion can be found in Kruschke [16], [22], [23]. Theoretical and empirical results regarding learnability, generalization, and network size can be found in, e.g., [24]-[26].

Given the desirability of small hidden layers, why not just build a network with an arbitrarily small number of nodes? First, for most learning applications it is not known what the minimal number of hidden nodes is. If one uses too few nodes, then the network has insufficient computational power to learn the training set to criterial accuracy, regardless of learning algorithm used. Second, back-propagation learning is slower, and more likely to encounter local minima or extensive plateaus on the error surface, when small hidden layers are used. For example, the minimal number of hidden nodes required to learn the XOR is two, but when only two nodes are built into the network, extended learning times are often encountered. In general, back-propagation learning is faster when larger hidden layers are used [10]. We try to satisfy these two competing constraints on hidden layer size by building a network with a

<sup>&</sup>lt;sup>1</sup>For non-infinitesimal changes, suppose that  $||\Delta w_i^s|| = p||w_i^s||$ , for some  $0 . Then we must have <math>\cos \theta < -p/2$  for the conclusion to hold. For small p, that condition is virtually the same as  $\cos \theta < 0$ . In other words, for non-infinitesimal changes, the conclusion of the lemma can be violated when the change vector  $\Delta w_i^s$  is at an angle just slightly obtuse to the weight vector  $(-p/2 \le \cos \theta < 0)$  or when it happens to be huge relative to the weight vector (p > 1). Empirically, such conditions occur so ephemerally as to be negligible.

<sup>&</sup>lt;sup>2</sup>There are also cases for which reducing the number of hidden nodes did not improve generalization (e.g., [21]). It is beyond the scope of this article to analyze in detail which particular applications will benefit from hidden layer bottlenecks. The premise is that at least some applications do benefit from bottlenecks.

large number of hidden nodes and dynamically reducing them as learning progresses. Our previous observations regarding the gain parameter suggest that a judicious manipulation of gains could accomplish this goal.

### Generalized Cost Function for Gain Competition

We can use the gain parameter to make the hidden nodes compete for the right to participate in the representation and in learning. The idea is to start with a large hidden layer of candidate nodes, and to impose a cost on nodes that are similar. In general, the cost of similar nodes can be expressed as

$$C = \sum_{i,j} h(g_i) h(g_j) s_{ij}$$
(12)

where *h* is a nondecreasing function of gain, and  $s_{ij}$  is some measure of the similarity of nodes *i* and *j*. The sum is taken over all nodes in the hidden layer. Thus if two nodes are similar and both have large gains, they make a large contribution to the cost. We can perform gradient descent on the cost *C* with respect to gain in order to decrease the gains of redundant nodes. (One might also perform gradient descent on *C* with respect to connection weights.) We desire the derivative to be locally computable in a network architecture compatible with back propagation. Thus, the gain of a node is adjusted simultaneously by gradient descent on error and by gradient descent on the cost *C* (12).

The Special Case of Gain Decay: One special case of the cost function defines h to be the identity function, and defines  $s_{ij}$  to be 1 if i = j and 0 otherwise. Then the cost function reduces to  $C = \sum_i g_i^2$ , and gradient descent on C with respect to  $g_i$  yields  $\Delta g_i \alpha - g_i$ . That is simply gain decay, analogous to weight decay, which has often been used in applications of back propagation (e.g., [18], [27]).

Another special case defines  $h(g_i) = g_i / (1 + g_i)$ , with  $s_{ij}$  again equal to 1 if i = j and 0 otherwise, so that  $C = \sum_i g_i^2 / (1 + g_i)^2$ . This is also a form of gain decay, in which the decay rate increases to a maximum of some small value of gain, and then decreases as gain gets larger. It is analogous to a cost function used by Rumelhart [19], [20], [28].

Those special cases of the cost function are particularly simple and easy to implement in a back propagation network. Unfortunately, gain decay alone does not take full advantage of the potential captured by the generalized cost function. We suggest now, and provide empirical evidence later, that gain decay excises slow-learning nodes rather than redundant nodes.

Specific Cost Function for Gain Competition: The measure of similarity used in the special cases of gain decay is really a degenerate case. We seek instead some measure of similarity which rises continuously from zero. Moreover, the measure must allow gradient descent on the cost of gains to be locally computable in a network compatible with back propagation. One such function is as follows:

$$s_{ij} = \frac{\left(\sum_{p} \operatorname{net}_{ip} \operatorname{net}_{jp}\right)^{2}}{\left(\sum_{p} \operatorname{net}_{ip}^{2}\right)\left(\sum_{p} \operatorname{net}_{jp}^{2}\right)}$$
(13)

where the sum is taken over all patterns, p, in a training sweep. The notation net<sub>ip</sub> denotes the net input to node *i* when pattern *p* is input. The similarity in (13) is a measure of the squared correlation of the net input values of nodes *i* and *j* across patterns. (It is not the usual definition of correlation because we have not subtracted the mean net input values.) It is always between zero and one, varies continuously, and it has a particularly simple derivative with respect to gain: zero. If we choose the identity function for h(g) in the generalized cost function *C*, then gradient descent on C with respect to gain yields the competition rule:

$$\Delta g_i = -\gamma \sum_{i} g_j s_{ij} \tag{14}$$

where  $\gamma \ge 0$  is a constant of proportionality, called the competition rate. A variant of (14) was first introduced by Kruschke [16], who used a slightly different definition for  $s_{ij}$ .

Local Computation: The value of  $\Delta g_i$  can be computed locally in a back propagation network if lateral connections between hidden layer nodes are allowed, that do not propagate activation or errors, but gain values. The strength on the lateral connection from node *i* to node *j* is  $s_{ij}$ , which can be computed with information available locally at the two nodes. Then  $\sum_j g_j s_{ij}$ is easily computed by summing across the lateral connections.<sup>3</sup>

### Simulation Procedure

The encoder problem [1] is an especially appropriate test for the gain competition scheme. In the encoder problem, a network with N input nodes and N output nodes is trained to map the N canonical basis vectors to themselves; i.e.,  $(1, 0, \dots) \rightarrow$  $(1, 0, \dots), (0, 1, \dots) \rightarrow (0, 1, \dots)$ , etc. "Good" solutions to the encoder problem use  $\log_2 N$  hidden nodes that together act as a binary encoder of the input. In that case, the hidden nodes are completely uncorrelated with each other. That is exactly the sort of situation that the gain competition scheme is supposed to encourage.

We used a network with eight input nodes, 16 hidden nodes and eight output nodes. To initialize the network, the weights on each node were chosen at random and then the weight vector of each node, including its bias weight, was normalized to Euclidean length one. The gain of each node was also initially set to one. Standard back propagation was performed; i.e., gradient descent on error with respect to weights, with a learning rate of 0.20 and a momentum of 0.85. Gradient descent on error with respect to gains was also performed on both the hidden and output layers, with the learning rate set to 0.20, and zero momentum. The learning was run until the maximum absolute error for every output node was less than 0.15.

Gain competition was simultaneously performed on the hidden layer. We wanted to impose as much competition as possible without extinguishing learning (error reduction), so we dynamically modulated the competition rate as learning progressed, in the following simple way. If learning was proceeding well, with the square root of the mean squared error (RMSE) falling rapidly, then competition was increased. If learning was slow, competition decreased. Specifically, the competition rate  $\gamma$ of (14) was a constant multiple of an exponential average of the change in RMSE over previous trials, divided by the RMSE of the present trial:

$$\gamma(\tau) = -\kappa \left( \sum_{t=0}^{\tau} \omega^{t} \Delta \text{RMSE}(\tau - t) \right) / \text{RMSE}(\tau) \quad (15)$$

where  $\gamma(\tau)$  is the competition rate on trial  $\tau$ , where  $\kappa$  is a non-negative constant of proportionality called the competition rate modulator, where the sum is taken over all previous trials t, and where  $\omega$  is a positive constant governing the weighting of the exponential average. In our simulations we set  $\omega = 0.70$ , and tested the competition rate modulator  $\kappa$  at several values between zero and two. Note that if the network is in fact learning, then  $\Delta$ RMSE should be negative, and so the competition rate  $\gamma$ 

<sup>3</sup>One can also easily derive  $\partial C / \partial w_{uc}$ , but computing the change in weight is not any simple variant of the feed-forward network for back propagation. One goal for future research is to construct a function  $s_{ij}$ , which yields gradient descent on C with respect to weights that is locally computable.

TABLE IV NUMBER OF PARTICIPATING HIDDEN NODES (NODES WITH NONZERO GAIN)<sup>a</sup>

	NON.	CERO GAIN)				
κ	0.0	0.5	1.0	2.0		
Mean	16.0	5.33	3.93	3.20		
Range	16-16	4-6	3-5	3-4		
SD	0	0.70	0.57	0.40		

<sup>a</sup>After learning with competition, for different values of  $\kappa$ .

should be positive. However, gradient descent algorithms like back-propagation can sometimes "overshoot" a local descent and cause  $\Delta$ RMSE to be greater than zero, consequently causing  $\gamma$  to be less than zero. To deal with that possibility, the value of the competition rate  $\gamma$  was actually set to the maximum of zero and the value computed in (15). Gain competition was not actually executed until the fifth sweep, so that the value of  $\gamma$ would reflect the smooth behavior of RMSE rather than the initial turbulence often seen in back propagation. Finally, gain was restricted to be non-negative so that competition could never "explode," driving some gains to large negative values and others to large positive values.

It was found that gain decay contributed almost nothing to the excision of hidden nodes (these results are reported later), and added significantly to learning time. We therefore did not include the diagonal terms of (12) in the computation of gain cost, C. Thus the change in gain due to competition was

$$\Delta g_i = -\gamma \sum_{j \neq i} g_j s_{ij} \tag{16}$$

(cf. (14)). Every factor in the right hand side of (16) is a non-negative number. Hence competition could only decrease the gains, never increase them.

#### Results

The results showed a large decrease in the number of participating hidden nodes as a function of the competition rate modulator,  $\kappa$ . Results are summarized in Table IV. Fifteen trials were run at each value of  $\kappa$ , each trial starting with a different set of random weights. (The same fifteen initializations were used for each of the values of  $\kappa$ .)

When there was no gain competition ( $\kappa = 0.0$ ), all 16 of the hidden nodes built into the network ended up participating in the representation. In fact, all 16 nodes had gains greater than 1 on every trial. From these results it is clear that back propagation alone does nothing to excise redundant nodes, but instead utilizes all the nodes available to it.

When only a small amount of competition was used ( $\kappa = 0.5$ ), the number of participating hidden nodes dropped to an average of only 5.33. All the other hidden nodes had gains of zero (to more than six decimal places). The number of participating hidden nodes did not vary much from trial to trial, as can be seen from Table IV.

As the gain competition was increased to  $\kappa = 2.0$ , usually (in 12 of 15 trials) only three hidden nodes remained. That is exactly the "good" solution, using  $\log_2 8$  hidden nodes, that the competition scheme was meant to encourage.

The benefits of gain competition do not come without some cost. As might be expected, learning time increased as the competition rate modulator  $\kappa$  increased. Table V shows that for  $\kappa = 2.0$ , learning time is almost three times that for  $\kappa = 0.0$ . It is important to notice that learning time is a negatively accelerating function of  $\kappa$ . Thus gain competition does not cause an explosive growth in learning time.

For comparison, an 8-3-8 network was tested on the encoder problem with fifteen random weight initializations, using the same procedure but without competition. The network reached

TABLE V Learning Time in Sweeps for Various Amounts of Competition

κ	0.0	0.5	1.0	2.0
Mean Range SD	218 105–303 55.3	403 203–722 129.9	497 274–769 162.3	600 353-824 145.0

TABLE VI EFFECTS OF GAIN DECAY ALONE			
к	0.5	1.0	
Mean hidden nodes	16	16	
Mean hidden gain	0.343	0.234	
Mean learning time	346	430	

criterion in an average of 328 sweeps, with a range of 195 to 455 sweeps, and a standard deviation of 66.8 sweeps. Thus adding competition takes less than twice as long as back propagation with gain. Since back propagation with gain was shown to be about twice as fast as standard back propagation, the competition scheme ends up taking no longer than standard back propagation.

The results reported so far have described the retirement of nodes from participation. On the other hand, if the competition rate modulator  $\kappa$  is set to much larger values, some interesting behavior occurs. The high competition initially suppresses all but one or two nodes. The network will reduce the error as best it can, but as the rate of error reduction slows, the competition rate  $\gamma$  gets smaller (by (15)). This allows new nodes to be recruited from the pool of retired nodes. One or more previously suppressed nodes grows to have positive gain, as a result of gradient descent on error with respect to gain. The newly recruited node is dissimilar to the previously participating nodes (otherwise competition would have continued to suppress it), and error can be reduced further (otherwise it would not have been recruited).

As suggested earlier, gain decay alone should not specifically excise redundant nodes, but only those that learn slowly. Here we present empirical support for that claim. Gain decay alone  $(\Delta g_i = -\gamma g_i)$  was applied using the same procedure as for competition without decay. Table VI shows the results for the same fifteen initializations as used in Tables IV and V, with  $\kappa$ equal to 0.5 and 1.0. In this context,  $\kappa$  is the "decay" rate modulator, rather than a "competition" rate modulator.<sup>4</sup> All the hidden nodes had gains greater than zero after learning; that is, none were excised at all. Gain decay had two significant effects: It increased the learning time; and, it reduced the mean hidden node gain. Clearly this form of gain decay is counterproductive, and so we have good evidence that competition is an improvement over decay alone.

#### Other Minimization Techniques

Several researchers have introduced other hidden layer minimization techniques, e.g., [16], [19], [20], [22], [23], [28]–[34]. No direct comparisons have been made of the relative performance of the alternative methods, in terms of learning speed and generalization improvement for particular applications. Each method introduces its own bias upon the type of representation learned by the network, and hence upon the generalization made by the network. For example, the gain competition method described here biases the network toward hidden layer representations that have uncorrelated node activations. That bias

<sup>4</sup>Not shown are results for  $\kappa = 2.0$ , which led to unstable oscillations even for very small learning step sizes.

might be appropriate for some applications, and not for others. Fortunately, the generalized cost function for gains (12) allows other definitions of similarity that might be more appropriate for specific applications.

In some applications it might be known in advance that specific kinds of bias are desired, and they might have nothing to do with minimization of hidden layers. For example, in visual pattern recognition, the system should generalize over spatial translations, and so the constraint of translational symmetry can be built in from the start [1]. Hidden layer minimization techniques, on the other hand, are usually intended for applications in which the appropriate domain-specific constraints are not known in advance.

### V. GENERAL DISCUSSION

#### Gain and Temperature

Both Boltzmann-type networks [18], [35] and back-propagation networks use sigmoidal activation functions. The "temperature" parameter used in Boltzmann machines can be thought of as the reciprocal of gain [6], [7]. In view of this apparent similarity it is important to point out the following differences.

Boltzmann machines are stochastic and activation values are binary. In these networks, temperature connotes stochasticity of the activation values. On the other hand, BPG is a deterministic network and therefore, gain (or its reciprocal, temperature) cannot connote randomness. But BPG activation functions may be interpreted as mean field approximations to binary stochastic nodes (e.g., [36]-[38]). The stochastic interpretation of temperature in Boltzmann machines corresponds to discriminating power in deterministic systems.

In Boltzmann and mean field networks, all nodes have the same temperature and are subject to a global "cooling" or "annealing" schedule (but cf. [39]). In Boltzmann machines, sufficiently slow cooling rates guarantee that the activation values for a given fixed set of weights will settle in a global energy minimum [40]. No corresponding claim is being made for BPG; there is no externally imposed annealing schedule for gains, and changing the gains by gradient descent does nothing to avoid local minima on the error surface.

### Gain and Natural Networks

There is evidence suggesting that the nervous system has mechanisms that modulate the neural response function in a manner similar to gain. Servan-Schreiber, Printz, and Cohen [41] proposed that a gain parameter helps explain the effect of biogenic monoamines, a type of neurotransmiter associated with neural responsiveness modulation. They used the gain parameter in back-propagation networks to simulate phenomena associated with catecholamine manipulations. There is also evidence that these amines play an important role in the modulation of learning [42].

# VI. CONCLUSION

We have introduced a gain parameter into the standard back propagation algorithm and shown it to be beneficial for speeding up learning and creating hidden layer bottlenecks for the goal of improved generalizations. It was shown that the gain of a node is a handle on its participation in representing the input, and on the node's ability to learn. Simulations confirmed that adaptive gains significantly accelerated standard back propagation. Other simulations showed that gain competition very effectively created hidden layer bottlenecks in the case of the encoder problem.

Not all applications will benefit equally from the techniques we have introduced. BPG speeds up learning when there are nodes which are learning successfully. When very few hidden nodes are included, there might be no successful nodes, and hence BPG might not significantly improve learning. The competition scheme is designed to favor solutions with uncorrelated hidden nodes. Our simulations showed that it works very well for the encoder problem, but if a "good" solution to some other particular problem uses highly correlated nodes, then the competition scheme might be counter-productive. The general cost function for gains allows other definitions of similarity that might solve this difficulty.

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# Motion Analysis of Multiple Moving Objects Using Hartley Transform

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Abstract —A new technique for motion estimation of moving objects is presented. The analysis of the Hartley spectrum indicates that the velocities of moving objects are related to the locations of the spectral peaks. The presented formulations and simulations demonstrate the applicability of this technique. This method is simple, fast, and computationally efficient.

#### I. INTRODUCTION

The estimation of the velocity of moving objects is required in many applications. Biomedical cell motion analysis, tracking dust storms and clouds, and industrial and military applications are a few important examples. Some researchers have used segment and match technique to acquire velocity information [1]-[3]. Static images are segmented and then feature points are matched to establish correspondence between objects in successive frames. This technique is sensitive to segmentation errors. The success of the algorithm is based on accurate segmentation of static frames, which is rarely satisfied in real world scenes, especially in a noisy environment.

Other researchers [4]–[6] have used differencing techniques to extract moving objects in a sequence. In their techniques two frames are subtracted and the resulting frame yields the difference between the two frames. This method has limitations as it requires 1) images to be exactly registered; 2) illumination to be invariant; and 3) the moving objects to be totally displaced. The temporal-spatial gradient techniques [7], [8] that utilize low-level estimation of frames, give best results only when the moving objects have smooth edges and the surfaces contain prominent texture. Their algorithm requires registered image sequences and the velocity estimate is approximate (it is close to accurate velocity at objects' boundaries, while this is not the case at the objects' surfaces). The above techniques have other limitations when multiple moving objects are in the sequence, the correspondence and occlusion problems are more severe.

Most researchers of time-varying image analysis use only two or three frames of a sequence. The analysis based on only a few frames does not take into account the complete information about the motion of objects. It has been shown [9], [10] that the human visual system requires an extended frame sequence in order to recover the structure of moving patterns. A longer sequence of frames allows the use of velocity information in analyzing the problem. Sethi and Jain [11] used a sequence of frames for finding the trajectories of feature points in an image sequence, and they used tokens to solve the correspondence problem, assuming smoothness of velocity. Selecting interesting

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