

LEARNING BY ERROR-DRIVEN DECOMPOSITION

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In this paper we describe a new selforganizing decomposition technique for learning high-dimensional mappings. Problem decomposition is performed in an error-driven manner, such that the resulting subtasks (patches) are equally well approximated. Our method combines an unsupervised learning scheme (Feature Maps [Koh84]) with a nonlinear approximator (Backpropagation [RHW86]). The resulting learning system is more stable and effective in changing environments than plain backpropagation and much more powerful than extended feature maps as proposed by [RS88, RMS89]. Extensions of our method give rise to active exploration strategies for autonomous agents facing unknown environments.

The appropriateness of our general purpose method will be demonstrated with an example from mathematical function approximation.

1 Introduction

Artificial neural networks have been found useful in widespread applications. One of the important problems concerning real-world applications is the slowness and instability of existing learning algorithms such as backpropagation [RHW86]. This slowness (which increases drastically with network size) is due to the fact, that

- all of the weights in the network are changed at once. Hidden units interact and are constantly facing changing information that delays the settling into useful hidden unit representations.
- the network architecture (number of hidden units, connectivity etc.) and parameter settings (e.g. learning rate, momentum) have to fit the problem, i.e. usually learning has to be repeated several times to find an appropriate network and parameter set.

Some approaches have already been proposed to eliminate the need for repeated trials with different architectures, such as the theoretical results on minimal hidden layer size [BH89] and constructive/destructive algorithms [FL90, Cha89].

In this paper we propose a radically different approach. Instead of constructing an architecture that fits a problem or function, we are decomposing the problem by splitting its input domain. Subdivision of input space is accomplished by an extended feature map. Associated with every feature map unit is a considerably small backpropagation (bp) network¹. It is used to approximate that part of the function, that is defined over the receptive field of its feature map unit. Instead of using equally defined patches or a decomposition that is just dependent on the probability distribution of the input space, we describe a new difficultness-sensitive decomposition method. Local approximation error of the bp-networks drives the distribution of feature map units. Thus an error-driven decomposition of the input domain is created. Difficult parts of the function yield fine grained patches (more bp-nets), while easier ones lead to a coarse grained decomposition and are approximated by less bp-networks.

Our approach can be seen as an application of the devide and conquer-principle (or problem decomposition) that has a long tradition in computer science [HS78] and artificial neural network learning [JJB90, MT90]. Its usefulness is widely accepted (e.g. finite element method [BP84] in numerical mathematics, patching [Far90] in geometric modeling). But traditional methods suffer from the fact that either the decomposition has to be given in advance or is performed in a strict, predetermined manner (e.g. splitting in halves). We introduce an error-driven technique to automatically derive a suitable task decomposition. Some considerable benefits will be demonstrated:

¹Actually just hidden units and their associated weight vectors are manipulated. Input and output units need not be duplicated.

- The learning process compared to backpropagation is sped up by using many small networks in a modular fashion rather than a single large one.
- Execution speed is enhanced by modularization, and it is possible to perform faster (gradient directed) search in input space, as it is used by a planning method described in [TML90, TML91].
- The problem of determining the number of hidden units is circumvented. The system performs an adaptive resource allocation in a way such that “difficult” parts of a function attract more networks (or hidden units) than easier ones. Destructive/constructive extensions can be incorporated easily.
- The instability (or relearning) problem is eased considerably, because additional patterns or slight changes of some old patterns only affect a small part of the whole network. Thus, their unstabilizing influence is locally restricted to a small number of hidden units.
- An active, error-driven exploration strategy can be derived with our method, if adaption to an unknown environment is desired.

We will proceede by introducing our novel learning technique and its architecture.

2 Function approximation as a learning process

In the sequel we will compare several connectionist approximation methods, whose approximation capabilities will be demonstrated by the mexican hat function $f(x, y) = (4(x^2 + y^2) - 2)e^{-\frac{(x^2+y^2)}{0.72}}$.

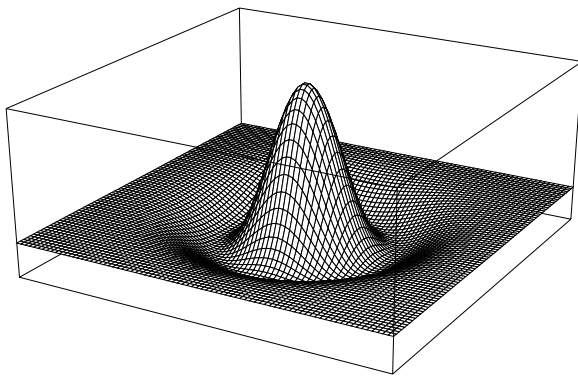


Figure 1: Mexican hat function

We start with approximation by a discrete step function using an extended feature map algorithm.

2.1 Approximation with a step function

In order to approximate a multidimensional function, we first introduce a technique which is related to feature maps [Koh84, RMS89]. In this approach the input space is devided into small regions, each of which is represented by a feature vector.

A feature map formally defines a step-function

$$FM : x \in I\!\!R^n \rightarrow p \in M$$

where M is a m -dimensional grid, i.e. the cartesian product of m indexsets I_i with $I_i \subset \mathbb{Z}$ and some neighborhood function. Every point in the input domain is assigned to some element of M . If we consider any element $m_k \in M$ a formal neuron, its receptive field is defined as

$$\mathcal{R}_k = \{x \in I\!\!R^n | FM(x) = m_k\}$$

The mapping FM is formed by adapting internal parameters $\{w_i \in I\!\!R^n; i \in M\}$ (called weights) by the following algorithm.

1. Generate/read (next) input $x \in I\!\!R^n$.
2. Determine node $j \in M$ whose parameter set (weights) is closest² to x .
3. Let $\mathcal{N}(j) \subset M$ be a symmetrically defined neighborhood of the winning node j in the m -dimensional grid. $\forall i \in \mathcal{N}(j)$ adapt the parameters according to the rule:

²This can be done by minimizing the Euclidean distance $\|w_j - x\| = \min_i \|w_i - x\|$, $i \in M$ or, in case of normalized weights by maximizing the inner product: $w_j \cdot x = \max_i \{w_i \cdot x\}$, $i \in M$

$$w_i(t+1) = w_i(t) + \alpha_i(t)(x(t) - w_i(t))$$

where $\alpha_i(t) = \epsilon(t)h_{ij}(t)$ consists of a stepsize $\epsilon(t)$ and a neighborhood term $h_{ij}(t)$ that is usually taken to be gaussian centered at the winning node j .

4. Change stepsize $\epsilon(t)$ and neighborhood function $h_{ij}(t)$, such that these are monotonically decreasing.
5. Goto step 1.

An extension of this algorithm was used by Ritter, Martinez and Schulten [RMS89] to perform function approximation. They assigned to every node in the FM-grid a set of output parameters that are adapted over time to minimize the LMS difference of the output value and the desired k -dimensional $f(x)$. Formally they introduced an output mapping OUT ³:

$$OUT : p \in M \rightarrow o \in I\!\!R^k$$

During learning any node of the feature map is assigned some output value. This value approaches the mean of the target values over its receptive field. This leads to a stepwise approximation of the function that is governed by the probability distribution of input space.

Just a small change to Kohonen's algorithm need to be given for this method: after step 3 an additional adaption step has to be introduced. It might, for example, be realized as a gradient descent on a quadratic error function (Widrow-Hoff or delta rule).

- 3.a $o_i(t+1) = o_i(t) + \alpha'_i(t)(f_i(t) - o_i(t))$ where $o_i(t)$ denotes the output weight vector of unit i at time t , $f_i(t)$ the i th dimension of the function to be approximated and $\alpha'_i(t)$ the stepsize at time t .

We used this algorithm to approximate the mexican hat function. Results obtained with a uniform distribution of input patterns are given in figure 2. After 850000 sample presentations learning converged. A plot of the distribution of feature map units and a plot of the approximated mexican hat function is shown.

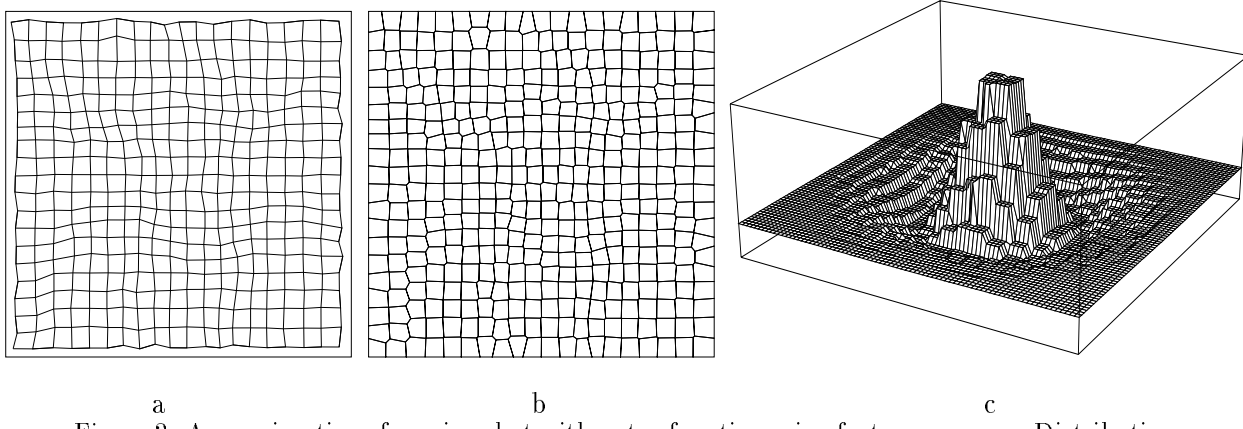


Figure 2: Approximation of mexican hat with a step function using feature maps. a. Distribution of units (20×20) b. Receptive fields c. Plot of approximation (tss=2.5).

Obviously this approximation is not optimal, i.e. with the same number of feature map units (patches) a better approximation can be derived (cf. figure 3.), if the size of receptive fields (subtasks) is allowed to change independently of the input distribution.

2.2 Error-driven decomposition

Obviously the above given example is not optimal with respect to the L_2 -norm. The overall error of the approximation, $E = \int_M P(x)|o(x) - f(x)|^2 dx$, is influenced by two factors:

- the probability distribution $P(x)$ of the input domain and
- the squared error (L_2 -norm) of the approximation

The contribution of a FM-unit i to this error is given by $E_i = \int_{R_i} P(x)|o(x) - f(x)|^2 dx$, i.e. the size of receptive fields should be determined not just by the probability distribution of the input space, but by the quality of the local approximation as well. We introduce an error-driven decomposition method, that leads to an even distribution of error over all patches.

To accomplish this goal, we extended the above algorithm by introducing into step 3. an error term

³This was enhanced by an approximation of the Jacobian to linearly interpolate between OUT -values. Their method relied on knowledge about the inverse of the function f .

$pss(t) = \sum_j (o_j(t) - f_j(t))^2$. On the long run it signals if some part of the problem is more difficult to approximate than others.

Thus we used the changed update rule:

$$w_i(t+1) = w_i(t) + \alpha_i(t)pss(t)(x(t) - w_i(t))$$

It leads to a stronger adaption step if the error is high than if approximation is already good.

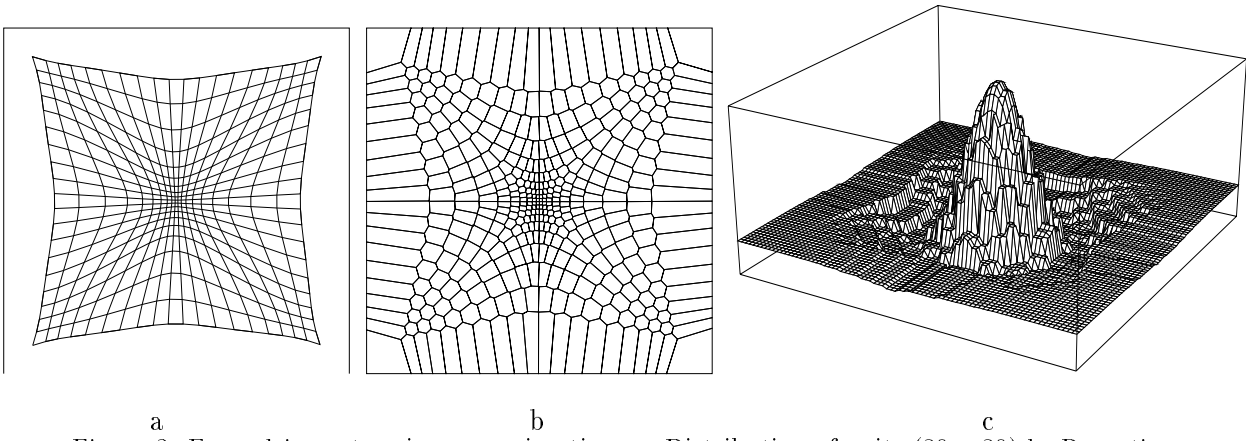


Figure 3: Error-driven stepwise approximation: a. Distribution of units (20×20) b. Receptive fields c. Plot of approximation ($tss=1.24$).

In figure 3 results after 900,000 presentations are shown using this update rule. Please note the distribution of units that is driven by the approximation error. A discussion of convergence properties of this method may be found in [Moe91].

So far we presented an error-driven decomposition method whose approximation capabilities are limited due to its stepwise manner. We will extend this to a piecewise linear and even nonlinear approximation.

3 Linear and nonlinear approximation

A straight forward nonlinear approximation can be tried with plain backpropagation networks⁴. Unfortunately a huge effort is needed just to learn the mexican hat function. We used a network with 100 hidden units to derive the result as given in figure 4. Training was performed on a 16k Connection Machine. After almost 1 billion of pattern presentations (batch training with 16k patterns) the run was stopped, since there was no further noticeable improvement. What should be noted is, that we in fact tried a dozen other architectures on Sun Sparc workstations without any better results.

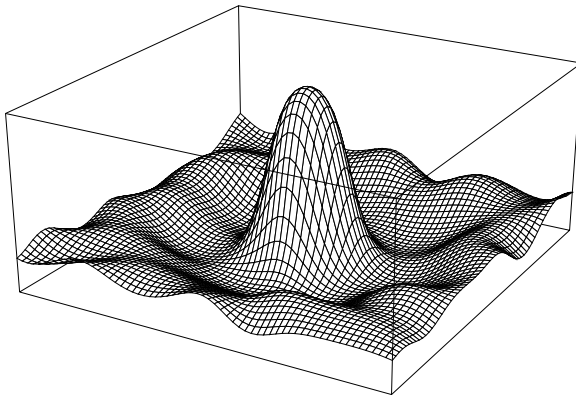


Figure 4: Approximation with pure Backpropagation ($tss=7.3$)

Since we are not really interested in approximating mexican hat functions with backpropagation, we omit a statistically valid investigation. But the slowness (almost one billion of pattern presentations!) of the pure bp-approach should be sufficiently clear.

⁴This is what we actually started with.

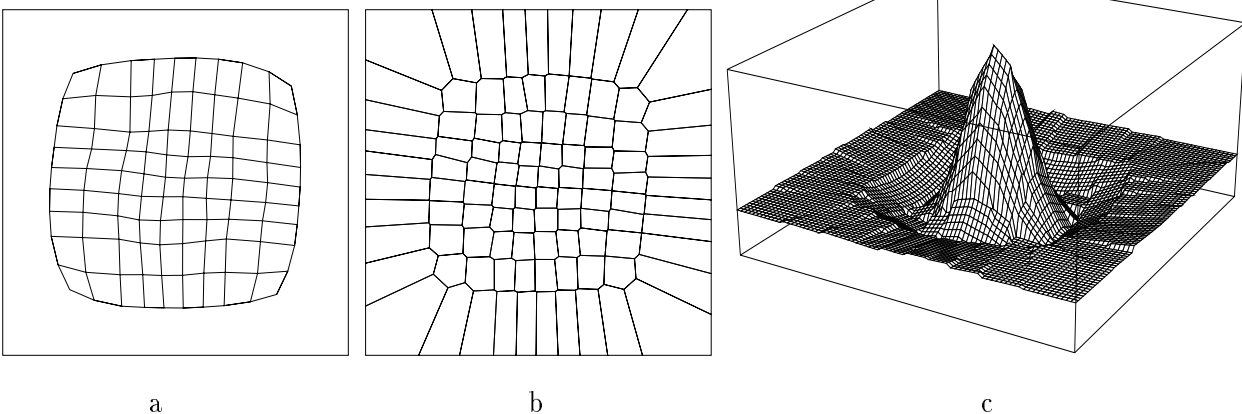


Figure 5: Linear approximation with decomposition (no hidden units). a. Distribution of linear nets (10×10) b. Receptive fields c. Plot of approximation ($tss=0.36$).

Our selfdecomposing algorithm, as described above, is now enhanced to allow for nonlinear approximations by using small backpropagation networks that are associated with feature map units. Each feature map unit is just used for gating the input to its associated network. Thus the receptive fields (as shown in fig. 5,6 b) determined by the feature map represent the input (or task) spaces of the simple bp-nets.

In our pseudo-algorithmic notion: step 3.a has to be exchanged by a backpropagation learning step.

- 3.b Train the backpropagation network associated to the winning node with the actual pattern pair $x, f(x)$. One training step!

This decomposition method has tremendous effects on backpropagation learning. With a 10 by 10 feature map, i.e. 100 small backpropagation nets, (in the above demonstrations we used 400 feature map units) the following results could be achieved:

- after 8 million pattern presentations the approximation without hidden units reached a quality as is shown in figure 5 ($tss=0.36$)
- results ($tss=0.07$) after 8 million pattern presentations are shown in figure 6. Networks with 3 hidden units were used.

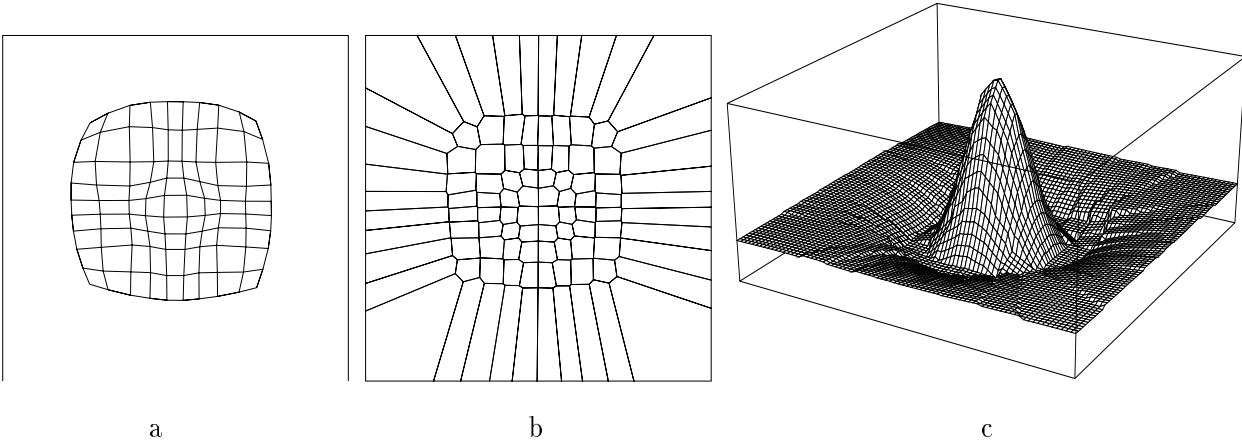


Figure 6: Nonlinear approximation with decomposition (3 hidden units). a. Distribution of backprop nets (10×10) b. Receptive fields c. Plot of approximation ($tss=0.07$).

The learning capabilities are quite stable with respect to parameter settings. We used a learning rate of .5 and a momentum term of .9 in those example runs. A survey of results is given in the following table, where the same parameter sets were used for all runs.

Method	Dimension	# hidden units per net	TSS	Patternpresentations in [millions]
Backprop	1	100	7.30	860
FM	20×20	0	2.52	1.0
FMirror-dvn	20×20	0	1.24	1.0
FMLinear	10×10	0	0.36	8.0
FMbackprop	10×10	3	0.07	8.0

4 Conclusion and future work

We described a nonlinear learning (or approximation) method that uses an error-driven decomposition technique. It combines an unsupervised learning method (feature maps) with a number of small supervised networks. The local adaption error of the networks (PSS) drives the learning of the feature map. The effect of feature map learning is an assignment of a receptive field to a bp-net in a way, that the variance of error over all FM-units is minimized. Usually a minimal overall error (with respect to the L_2 -norm) is approached.

It should be noted however, that there is no general convergence proof for our method. But during our extensive testing of the method we never observed divergence. In the future we will try to generate a case where this behavior can be provoked. Other extensions, that are being pursued, include:

- smoothness at the receptive field borders needs to be enforced. We are currently investigating how to combine gradient informations in neighboring networks such that the resulting patches are C_0 or even C_1 .
- overlaying of multiple grids with different bp-architectures competing for larger receptive fields.

To summarize: The motivation behind our approach was the need for a learning method that allows

- an easy coarse grained parallelization e.g. on transputer nets,
- faster convergence when learning real-world problems such as robot kinematics and dynamics. This is accomplished by the use of many simple networks (with few hidden units) instead of one big network, since in the latter case hidden unit representations are difficult to develop.
- active exploration, i.e. the focus of attention on those parts of workspace that are important at the moment and whose approximations are still not satisfying. It is straight forward how our learning method can be used to generate input for those areas where knowledge is still inadequate [TM91].
- easier relearning, i.e. in case that the learning system is facing a constantly changing environment fast relearning is necessary. Unfortunately pure backpropagation is not well suited for online training with open training sets. Our decomposition allows to keep certain changes local.

In this paper we presented a first demonstration of the capabilities of our new technique. Scaling to real world applications in the robotics domain are under investigation.

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