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Nonlinear Autoassociation Is Not Equivalent to PCA

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A common misperception within the neural network community is that even with nonlinearities in their hidden layer, autoassociators trained with backpropagation are equivalent to linear methods such as principal component analysis (PCA). Our purpose is to demonstrate that nonlinear autoassociators actually behave differently from linear methods and that they can outperform these methods when used for latent extraction, projection, and classification. While linear autoassociators emulate PCA, and thus exhibit a flat or unimodal reconstruction error surface, autoassociators with nonlinearities in their hidden layer learn domains by building error reconstruction surfaces that, depending on the task, contain *multiple* local valleys. This interpolation bias allows nonlinear autoassociators to represent appropriate classifications of nonlinear multimodal domains, in contrast to linear autoassociators, which are inappropriate for such tasks. In fact, autoassociators with hidden unit nonlinearities can be shown to perform nonlinear classification and nonlinear recognition.

1 Introduction ---

An autoassociator is a feedforward connectionist device whose goal is to reconstruct the input at the output layer. When used with a hidden layer smaller than the input or output layer and linear activations only, the autoassociator implements a compression scheme that was shown to be equivalent to principal component analysis (PCA)—also known as *singular value decomposition*—by Baldi and Hornik (1989). Another interesting and related result was obtained by Bourlard and Kamp (1988), who claim, “For

autoassociation with linear output units, the optimal weight values can be derived by standard linear algebra, consisting essentially in singular value decomposition (SVD) and making thus the nonlinear functions at the hidden layer completely unnecessary." They backed their claim using theoretical considerations, while Cottrell and Munro (1988) arrived at the same conclusion independently, using an experimental methodology.

Paradoxically, nonlinear autoassociators are also famous for their capacity to solve the encoder problem (Rumelhart, Hinton, & Williams, 1986), a problem that cannot be solved by PCA because of the singularity of the principal components.¹ Moreover, recent research suggests that nonlinear autoassociators are well suited for classification of certain types of nonlinearly separable and multimodal domains (Japkowicz, Myers, & Gluck, 1995; Petsche et al., 1996), another task for which PCA and linear autoassociation are not appropriate because of their linear restriction.

In spite of the obvious contradiction exhibited by these facts, there has been little effort at explaining the source of this inconsistency. The purpose of this article is to demonstrate experimentally that Bourlard and Kamp's (1988) claim does not necessarily hold and to analyze the differences between PCA and nonlinear autoassociators when it does not. In particular, we test the performance of nonlinear autoassociators relative to PCA and linear autoassociation on a nonlinear multimodal classification problem inspired by the projected topology of a real-world domain. After showing that there is indeed a difference in the classification accuracy obtained by the linear and nonlinear devices on this domain, we explain this difference by comparing the reconstruction error surfaces constructed by each system over the test domain.

More specifically, our experiments demonstrate that nonlinear sigmoidal single-hidden-layer autoassociators may sometimes interpolate sets of points differently from PCA. In particular, we show that on the test domain considered, while PCA (or equivalently, linear autoassociation) exhibits flat or unimodal reconstruction error surfaces, nonlinear sigmoidal autoassociators build error reconstruction surfaces that contain multiple local valleys. In fact, the reconstruction error surfaces built by the sigmoidal autoassociators are closely related to the reconstruction error surface built by a radial basis function autoassociator, thus suggesting that when the sigmoidal autoassociator does not operate like PCA—computing a globalized solu-

¹ Kruglyak (1990) and Phatak, Choi, and Koren (1993) demonstrate that for any N , the N -input, N -output encoder problem can, actually, be *represented* by an autoassociator of only two nonlinear hidden units. The fact that large encodings can actually be *learned* by two-hidden-unit autoassociators was demonstrated by Zipser (1989), who showed that such devices are capable of learning how to encode the position of a spot (or cluster) of normally distributed points appearing at random locations on a 10×10 squared array.

tion to the interpolation problem—then it operates like a radial basis function autoassociator—computing a localized solution to the interpolation problem.²

We begin our study by discussing the limitations of Bourlard and Kamp's (1988) claim. We then demonstrate experimentally that these limitations can indeed be exceeded by standard problems within the context of classification and we analyze these results, concluding with some speculations as to why sigmoidal autoassociators sometimes do and sometimes do not operate like PCA.

2 Limitation of the Conventional Wisdom

When considering the results of Baldi and Hornik (1989), Bourlard and Kamp (1988), and Cottrell and Munro (1988) and the evidence that nonlinear autoassociators can solve problems that cannot be solved by PCA (Rumelhart et al., 1986; Japkowicz et al., 1995; Petsche et al., 1996), an important question comes to mind: Does the Bourlard and Kamp (1988) claim hold in all cases, or does it depend on certain assumptions concerning the underlying domain or the autoassociator?

A careful look at the discussion laid out in Bourlard and Kamp (1988) reveals that the claim actually does depend on a particular condition. More specifically, let F be the nonlinear function present at the output of the hidden units of a nonlinear autoassociator (F is the function that makes the autoassociator nonlinear). The assumption Bourlard and Kamp (1988) make in order to carry out their demonstration is that for small values of x , $F(x)$ can be approximated as closely as desired by the linear part of its power series expansion. However, this means that $x = h$, the vector of presynaptic hidden unit activations (i.e., the hidden unit activations prior to their transformation by F), must be in the linear range of the squashing function. This suggests that nonlinear autoassociators do not necessarily emulate PCA when the net inputs are outside the linear range of the squashing function.

The remainder of the article demonstrates experimentally that there are indeed differences between the linear and nonlinear schemes and that these differences have practical consequences for the task of classification.

3 Experiments

We conducted two sets of experiments for this article.

² Although both the sigmoidal and radial basis function autoassociators are nonlinear, all further references to nonlinear autoassociators correspond to references to sigmoidal autoassociators.

3.1 Specification of the Devices and Description of the Task.

3.1.1 Devices. The devices we used in our experiments are PCA and three autoassociators. From an analytical point of view, the four devices differ: two of them are purely linear, and the other two have nonlinear capabilities. The three autoassociators are connectionist methods that differ from one another with regard to the type of hidden and output layer they use. In more detail, we compare a one-hidden-layer autoassociator whose hidden and output layers are both linear (L-L); a one-hidden-layered autoassociator whose hidden layer is nonlinear but whose output layer is linear (NL-L); and a one-hidden-layered autoassociator whose hidden layer and output layer are both nonlinear (NL-NL). In each of these devices, the function used in the nonlinear units is the usual logistic function.

3.1.2 Task. The operation of the four devices was contrasted on the task of classification. A formal definition of the classification problem typically involves an input vector x and a discrete response vector y , which are such that the pair (x, y) belongs to some unknown joint probability distribution, P . The goal of classification is to induce a function $f(x)$ from a set $(x_1, y_1), (x_2, y_2), \dots, (x_N, y_N)$ of training examples, so that $f(x)$ predicts y . Here we assume that y is a binary vector. Although classification by neural networks is typically performed using a discrimination network,³ it can also be performed using a recognition approach implemented by PCA or the autoassociator. This approach is discussed in Oja (1983) for PCA and in Hanson and Kegl (1987) for autoassociators and operates as follows.⁴ During the training phase, the autoassociator is taught how to reconstruct examples of the concept. Once training is completed, classification is performed on a new vector x_{Test} by comparing its reconstruction error⁵ to a threshold, and assigning it to the conceptual class if the reconstruction error is smaller than this threshold and to the other class otherwise. The idea behind this recognition-based classification scheme is that since the autoassociator is trained to compress and decompress examples of the conceptual class only,

³ Using such a scheme, y_i , the response variable associated with input vector x_i , takes on (in the simplest case) a value of 1 or 0, which is interpreted to mean that x_i belongs or does not belong to the conceptual class of the problem. After training the network to approximate the function $f(x)$ from which the training examples are believed to have been generated, it is expected that if the training set was appropriately designed, the network will be able to compute the appropriate label (1 or 0) for any input vector of size d , the size of the input layer, even if that vector did not appear in the training set.

⁴ We describe the method for autoassociators, taking into consideration that it is similar for PCA.

⁵ The reconstruction error corresponds to $\sum_{j=1}^d [x_{Test}^j - g(x_{Test}^j)]^2$, where g is the function implemented by the autoassociator and x_{Test}^j and $f^j(x_{Test})$ represent input unit j and output unit j of the network, respectively.

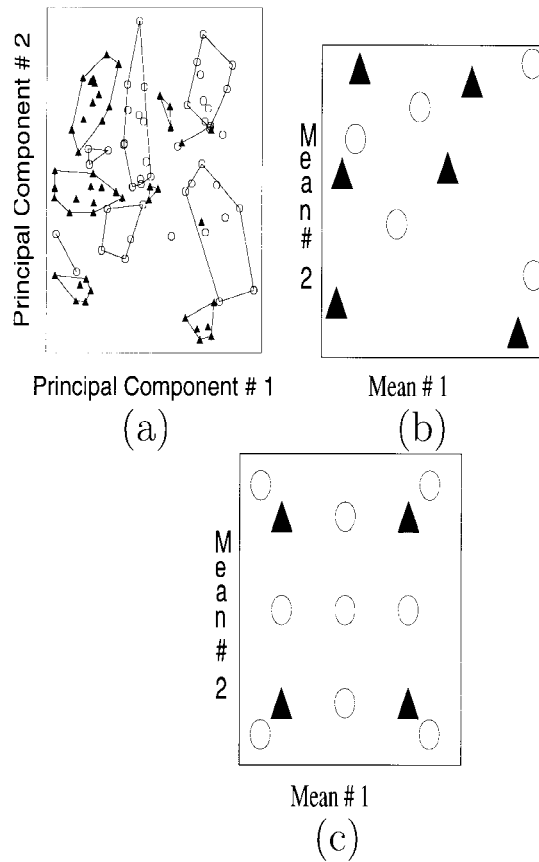


Figure 1: Transition from the sonar detection domain to the artificial nonlinear domain.

when tested on a novel data point, it will compress and decompress it appropriately if this example belongs to the conceptual class, but it will not do so appropriately if the example does not belong to the conceptual class.

3.2 Specification of the Test Domain. In order to compare and explain the classification performance of the four devices considered in this study, two experiments were conducted for each classifier on a two-dimensional artificial domain. This problem was inspired from real-world data as shown in Figure 1, which illustrates the transition from real-world to artificial data. Specifically, Figure 1a displays a two-dimensional representation of sonar target recognition data available from the University of California at Irvine Repository for Machine Learning. These data were compressed from 60

features to 2 using PCA. In this plot, black triangles and white circles correspond to actual points (mines and rocks, respectively), and data clusters are indicated by polygons. Figure 1b shows the same two-dimensional representation of sonar target recognition data, but this time with means replacing the clusters.⁶ Although this figure is just a two-dimensional projection of the original data set, it clearly indicates the typical kind of problem that the autoassociator has to deal with: multimodality. We chose a simple two-dimensional abstraction of the multimodality problem for benchmarking purposes and explaining the capabilities of autoassociators. This artificial domain, illustrated in Figure 1c, consists of four conceptual data clusters with means represented by black triangles and nine counterconceptual data clusters with means represented by white circles. In more detail, the conceptual clusters are located at points (.2, .2), (.2, .8), (.8, .2), and (.8, .8), respectively; the counterconceptual clusters are located at points (.1, .1), (.1, .9), (.2, .5), (.5, .2), (.5, .5), (.5, .8), (.8, .5), (.9, .1), and (.9, .9), respectively. Each cluster is composed of 50 points normally distributed around these means and with variance $\sigma^2 = .01$. The counterconceptual clusters are further divided into internal and external clusters. Internal counterconceptual clusters correspond to the counterconceptual data that are located inside the convex hull defined by the conceptual data, while external counterconceptual clusters correspond to the counter-conceptual data located outside this convex hull.

3.3 Experiment Set 1. The first experiment we conducted consisted of comparing the classification accuracy of the four systems on the test domain of Figure 1c. After having been tuned to their optimal settings, the four devices were trained on conceptual data, and a threshold was established by fitting the reconstruction error obtained on additional conceptual data to a gaussian and setting a boundary corresponding to a prespecified confidence interval. Subsequently, the systems were tested on a testing set containing both instances and counterinstances of the concept.

3.3.1 Tuning the Devices. The number of hidden units that the nonlinear connectionist systems used was determined by running each network with 1, 2, 4, 8, 16, 32, and 64 hidden units on five cross-validation sets containing 25 points per cluster. It was concluded that in order to reach a good classification performance, the two nonlinear autoassociators had to be trained with 16 hidden units. It was also shown that the networks had converged by Epoch 2000, which was thus selected as a stopping point. Optimization in this experiment was performed using the backpropagation procedure with standard learning rate and momentum of 0.05 and 0.9, respectively. For PCA, since the test domain is two-dimensional, only one or

⁶ Clusters of fewer than three points were ignored.

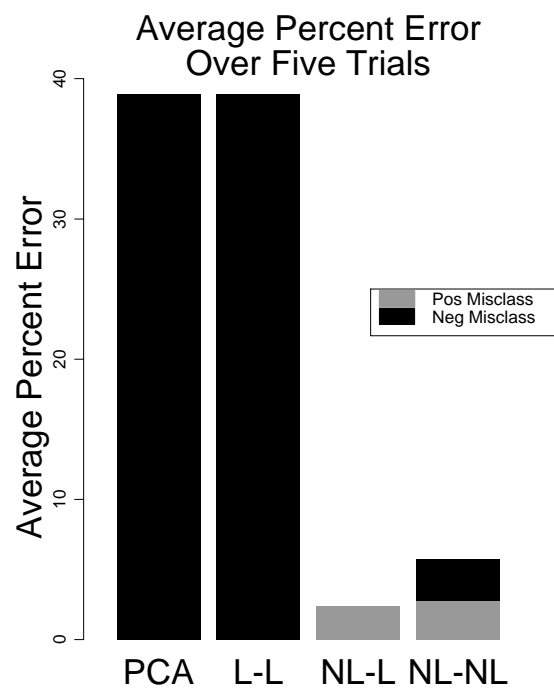


Figure 2: Classification error rate of the four classification schemes. Positive misclassification rates are indicated in gray; negative ones are indicated in black.

two principal components could be used. It was determined that one principal component yields a better classification rate than two, and therefore the PCA experiment was conducted using a single principal component. Linear autoassociation performed better as well with a single hidden unit than with two or more; therefore, it too was tested with a single hidden unit. The stopping criterion, learning rate, and momentum that the linear autoassociator used were the same as those used by the nonlinear devices. In the four systems, the threshold was set so as to allow for a 97% confidence interval.

3.3.2 Accuracy Rates. The results we obtained in this experiment are presented in Figure 2. This figure plots the average classification error rates over five trials obtained by the four systems on the test domain of Figure 1c. The graph shows that PCA and L-L obtained very large classification error rates, whereas the two nonlinear autoassociators obtained low error rates. This demonstrates that while PCA and L-L can technically be used for classification, they are not that useful on the problem of Figure 1c, whereas both

NL-L and NL-NL yield acceptable classification rates on that problem.⁷ This result thus confirms that there can be a difference in the computations performed by PCA (or linear autoassociation) and nonlinear autoassociation and that this difference has practical consequences for the task of classification. In order to analyze this difference in more depth, we conducted further experiments.

3.4 Experiment Set 2. These experiments sought to explain the results obtained in the previous set of experiments by computing and plotting the reconstruction error surfaces constructed by PCA and the three connectionist systems after being trained on the conceptual data (the data summarized by black triangles in Figure 1c). In other words, we were interested in finding out how the interpolation strategy that the various devices considered in this study used differ from one another.

3.4.1 Tuning the Devices. For experiment set 2, two different hidden unit settings were tested: expansion and compression. This was done so as to find out whether the systems operate qualitatively differently when used in a nonbottleneck or in a bottleneck fashion. In the expansion setting, the same 16-hidden-unit setting was used for the nonlinear systems as those used in the first experiment, while PCA and the linear autoassociator were tested with two principal components or hidden units. In other words, the optimal nonbottleneck setting for the nonlinear systems was compared to the only nonbottleneck setting possible for the linear systems.⁸ All the other parameters remained the same as in experiment set 1, except for the learning rate of L-L, which had to be increased from 0.05 to 0.1 for full convergence to take place. In the compression setting, the number of principal components and hidden units were restricted to one for all the systems since it is the only bottleneck setting available given that the test domain is two-dimensional. All the other parameters remained the same as in experiment 1.

3.4.2 Expansion Results. The results we obtained using the expansion setting of experiment set 2 are presented in Figure 3, which displays three-dimensional plots of the error ratio surfaces constructed by NL-L and NL-NL, respectively. The results for PCA and linear autoassociation are not displayed since they exhibit a flat reconstruction error surface. In the graphs of Figures 3a and 3b, the plots are drawn along the x -, y -, and z -axes where x corresponds to the x -axis of the input domain, y to its y -axis, and z is the

⁷ Nonlinear autoassociation-based classification was previously used in practical settings involving a CH-46 helicopter gearbox and motor monitoring (Japkowicz et al., 1995; Petsche et al., 1996) and yielded accurate results.

⁸ For the linear autoassociator, hidden layers greater than two are actually possible, but they are equivalent to hidden layers of size two.

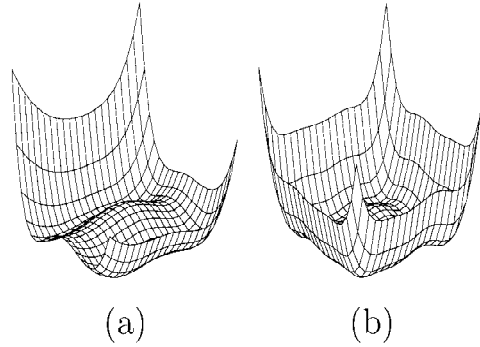


Figure 3: (a) Reconstruction errors of NL-L and (b) NL-NL, with 16 hidden units on the artificial domain generated from Figure 1c.

error ratio at every point considered in the space such that:

$$z(x, y) = \frac{1}{\gamma}((A(x) - x)^2 + (A(y) - y)^2). \quad (3.1)$$

In the formula, x and y represent the two dimensions of each data point, $A()$ is the function realized by the trained autoassociators, and γ is a model-dependent scale factor used for plotting purposes. The plots of Figures 3a and 3b are particularly helpful for understanding the nature of the solutions computed by the nonlinear autoassociators and contrasting them to the solution obtained by the linear autoassociator and PCA. In particular, they show that these error surfaces are qualitatively different from the ones computed by PCA and the linear autoassociator since, while the nonlinear autoassociators build multiple-local-valley representations of the underlying domain (see Figures 3a and 3b), PCA and the linear autoassociator learn how to reconstruct the domain perfectly and exhibit a flat reconstruction error surface. This suggests that the solutions computed by autoassociators with nonlinearities in their hidden layer use a multimodal interpolation bias that the linear methods do not use.⁹ Our results of Figure 3 can be used to ex-

⁹ As expected from Baldi and Hornik (1989), our results show that the linear autoassociator is capable of emulating PCA. However, in order for the two paradigms to be equivalent, the linear autoassociator had to be trained with a learning rate of 0.1 (instead of the learning rate of 0.05 used by the nonlinear systems). When a learning rate of 0.05 was used, the linear autoassociator got stuck in a saddle point (see Baldi & Hornik, 1989) and returned a constant output corresponding to the mean of the four training clusters. This observation is interesting in its own right because it underlines the difference between linear and nonlinear autoassociators even prior to the linear autoassociator's full convergence. Indeed, it suggests that the linear autoassociator tackles the reconstruction problem globally, using a unimodal bias, whereas the nonlinear autoassociators use local

plain the nonlinear autoassociators (NL-L and NL-NL) results of Figure 2, which show that nonlinear autoassociators are capable of classifying the nonlinearly separable and multimodal domain of Figure 1c. Indeed, since the reconstruction error surfaces of the nonlinear autoassociators include several local valleys centered at each of the positive clusters (see Figures 3a and 3b), the error ratios of the counterconceptual data can all be appropriately high relative to their conceptual counterparts, whether they are located in the interior, the sides, or the exterior of the square underlying the conceptual components. Thus, all the counterconceptual data get appropriately differentiated with respect to the conceptual data. The classification results obtained by PCA and linear autoassociation in Figure 2 will be explained next, since they were obtained with bottleneck devices.

3.4.3 Compression Results. The results obtained in the previous set of experiments demonstrate that nonlinear autoassociators are not equivalent to linear autoassociators or PCA in the case where the number of hidden units exceeds the number of input units or when the number of principal components is equal to the domain dimensionality. Autoassociators, however, are most typically used as compression devices with a number of hidden units smaller than the number of input units. We now discuss whether the same result also holds in this case. Figures 4a, 4b, and 4c display the reconstruction error surfaces obtained by PCA or L-L (Figure 4a), NL-L (Figure 4b), and NL-NL (Figure 4c) using a single principal component or hidden unit. These reconstruction errors were also obtained using equation (3.1). The reconstruction error surfaces obtained by these devices show that none is appropriate for full classification of the test domain of Figure 1c since they can only classify all (or most) of the data (whether conceptual or counterconceptual) in one diagonal of the input space as conceptual and all (or most) of the other data (whether conceptual or counterconceptual, again) as counterconceptual. In other words, because they do not interpolate the training set fully, these devices are not useful for classification, and this explains the high classification error rate obtained by the linear systems in Figure 2.¹⁰ Nevertheless, these results are interesting since, as in the non-bottleneck case, they do illustrate the difference between the linear and the nonlinear schemes. Indeed, while the linear systems display an undistorted surface with no saddle points (see Figure 4a), the nonlinear ones have saddle

strategies characterized by their multimodal interpolation biases.

¹⁰ While for the test domain used in this study, classification can be achieved only if using a number of hidden units greater than the number of input units, in the work of (Japkowicz et al. (1995) and Petsche et al. (1996), the number of hidden units is smaller than the number of input units. This suggests that the hidden-to-input unit ratio does not have any bearing on the behavior of nonlinear autoassociators in classification tasks, since what really matters is their capacity to interpolate the training set regardless of how many hidden units that may take.

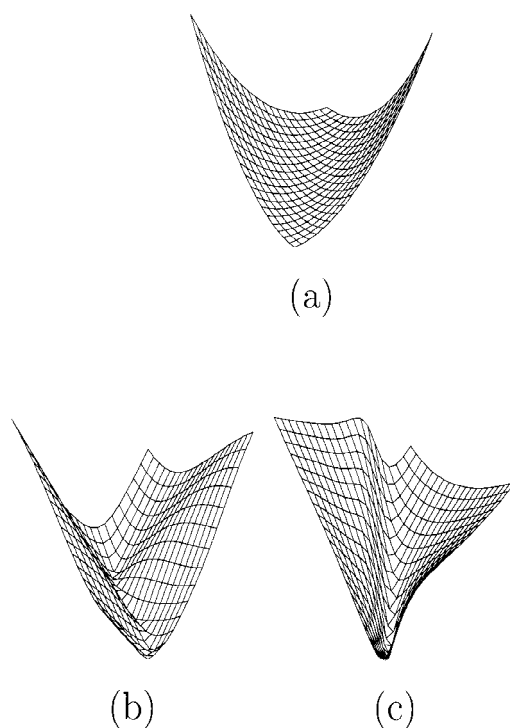


Figure 4: Reconstruction errors of (a) PCA or L-L (with one principal component or one hidden unit), (b) NL-L, and (c) NL-NL (with one hidden unit) on the artificial domain generated from Figure 1c.

points (see Figures 4b and 4c). Thus, the compression results demonstrate that nonlinear autoassociation is not equivalent to linear autoassociation or PCA even when the number of hidden units or principal components is smaller than the number of input units.

4 Discussion

We now analyze the results we obtained and explain why our observations depart from the expectations derived from the past literature. In addition, we compare the results of the nonlinear autoassociators to the result of another well-known paradigm, radial basis functions, and speculate on what nonlinear autoassociators compute and why they do so.

4.1 Nonlinear Autoassociation Versus PCA and Linear Autoassociation. The difference between the linear and nonlinear systems can be ex-

plained by the fact that for the particular test domain considered,¹¹ Bourlard and Kamp's (1988) assumption has been violated. Indeed, their discussion was shown to hold only in the case where the inputs to the nonlinear activation function of the hidden units remain in the linear range of the squashing function. An observation of the presynaptic activations of the 16 hidden units of NL-NL reveals that, on average, these values are equal to $\mu = 0.6953$ with variance $\sigma^2 = 0.1919$. Similarly, for NL-L, we found that $\mu = .3222$ and $\sigma^2 = .0548$. Although for input values in the $[-1, 1]$ interval, the sigmoid function is close to linear, our results seem to suggest that a small violation of the Bourlard and Kamp (1988) assumption can make a big difference in certain contexts. While this difference might be marginal and thus overlooked when considering the hidden layer alone, it is magnified in the output layer, as suggested by the plots of Figures 3a and 3b, which do not display a flat surface of the sort computed by PCA or autoassociation. This remark also holds for the encoder problem of Rumelhart et al. (1986), which can be solved by an autoassociator with nonlinear hidden units but not by a linear one. In this problem, nonlinearities in the hidden layer are also shown to have an impact.

4.2 What Do Nonlinear Autoassociators Compute? A Comparison with RBF Autoassociators. We will now attempt to establish the nature of the computations taking place in the nonlinear autoassociators by comparing their reconstruction error surfaces to the reconstruction error surface obtained by a radial basis function (RBF) autoassociator with gaussian activations in the hidden units and linear output activations. The reconstruction error surface obtained by such a device on the domain of Figure 1c is plotted in Figure 5. The particular RBF network that yielded this figure has a capacity of four hidden units, and the variance of the gaussian functions is fixed at and set to 2.8. The reconstruction error surface of this autoassociator was computed in the same fashion as that of the other autoassociators, using equation (3.1). The similarity between the reconstruction obtained by the nonbottleneck, nonlinear autoassociators (especially NL-L, which also has linear output activations) and the RBF autoassociator suggests that, like the RBF network, the nonlinear autoassociators use their 16 hidden units to put centers down over the four clusters representing the training data, and evaluate the distance of the testing data to these centers.¹²

This result, together with the results of Bourlard and Kamp (1988), suggests that nonlinear autoassociators can resort to one of two modes of operation: either they operate like PCA, seeking a globalized solution to the reconstruction problem they are trying to solve, or they operate like an RBF autoassociator, seeking a localized solution.

¹¹ For other classes of domains, this might not be the case.

¹² We thank Gary Cottrell for suggesting this comparison.

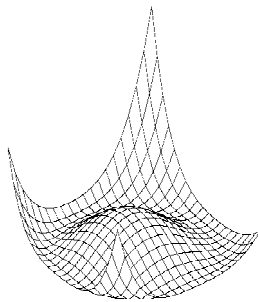


Figure 5: Reconstruction errors of the RBF autoassociator on the artificial domain generated from Figure 1c. The network has a capacity of four hidden units, and the variance of its gaussian functions is fixed and set at 2.8.

4.3 On the Duality of Nonlinear Autoassociators: Our Speculation.

The question we now ask is: When does a sigmoidal autoassociator resort to one or another of the two modes of action described in the previous section? Because of the current lack of technology available for analyzing feedforward neural networks fully, we can only speculate as to when each phenomenon will take place.

In particular, we suggest that the sigmoidal autoassociator will make use of its nonlinearities when the data set it attempts to reconstruct is multimodal and the different clusters that compose it are very spread out.¹³ Indeed, if the data set is unimodal or if it is multimodal but can easily be confused for unimodal data, then every point in the training set activates hidden unit values located in the same vicinity that increase or decrease monotonically. Such data can thus be approximated linearly, in the same fashion as they would be by a linear autoassociator. On the other hand, if the data set is multimodal and the different clusters constituting it are very spread out, then approximation of their hidden unit activation values by a linear function will fail because of the discontinuities introduced by the multimodal nature of the domain. In such cases, the sigmoidal part of the activation function is very useful since it permits the local processing of separate clusters in the same fashion as RBF autoassociators. Indeed, by using a sigmoidal function, a hidden unit can map all the data contained in different clusters to approximately the same default activation value (of 0 or 1) while it can map the data in some other clusters, which fall in the linear part of the sigmoidal function, to more interesting values. Each hidden unit can then use the same strategy for different clusters, and thus a

¹³ Support for this speculation can be found in Japkowicz (1999), which shows a clear decrease in classification accuracy by NL-NL as the conceptual clusters of the domain in Figure 1c are moved closer to one another.

global solution can be found by the overall network by compounding the localized solution computed by each hidden unit.

Because in the encoder problem, the autoassociator is expected to reconstruct data points located far away from each other—since they are located at extreme corners of the input space—this speculation also explains why the nonlinear autoassociator resorts to a localized solution not available to PCA in that case.

5 Summary

The purpose of this article was to address the paradox raised by the claim of Bourlard and Kamp (1988) that, in autoassociation, nonlinearities in the hidden units are completely unnecessary, given that nonlinear autoassociators are known to perform tasks that cannot be solved by PCA or linear autoassociation. We began by demonstrating that although both PCA (or linear autoassociation) and nonlinear sigmoidal autoassociation can be used for classification, the linear systems are not appropriate for certain types of multimodal and nonlinear domains, whereas the nonlinear systems are capable of classifying such domains accurately. An explanation of this result is provided by the observation of the interpolation surfaces constructed by the linear and nonlinear systems when trained on the conceptual examples of our classification test domain.

The difference in the interpolation and, as a result, the classification scheme used by the linear and nonlinear devices thus contradicts the claim by Bourlard and Kamp (1988) initially considered. This contradiction is resolved by extracting the assumption on which the claim is based and showing that this assumption is not necessarily always verified.

In a last step, we note that when the nonlinear autoassociator does not operate in a linear fashion, it emulates an RBF autoassociator and thus computes a localized solution to the problem it attempts to solve. Speculations as to when nonlinear autoassociators do resort to a linear operative mode and when they do not do so are then formulated and conclude the article.

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