

Neural network modeling of memory deterioration in Alzheimer's disease

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Abstract

The clinical course of Alzheimer's disease (AD) is generally characterized by progressive gradual deterioration, although large clinical variability exists. Motivated by the recent quantitative reports of synaptic changes in AD, we use a neural network model to investigate how the interplay between synaptic deletion and compensation determines the pattern of memory deterioration, a clinical hallmark of AD. Within the model we show that the deterioration of memory retrieval due to synaptic deletion can be much delayed by multiplying all the remaining synaptic weights by a common factor which keeps the average input to each neuron at the same level. This parallels the experimental observation that the total synaptic area per unit volume (TSA) is initially preserved when synaptic deletion occurs. By using different dependencies of the compensatory factor on the amount of synaptic deletion one can define various compensation strategies which can account for the observed variation in the severity and progression rate of AD.

1 Introduction

Alzheimer’s disease (AD) is the major degenerative disease of the brain, responsible for a progressive deterioration of the patient’s cognitive and motor function, with a grave prognosis (Adams and Victor 1989). Its clinical course is usually characterized by gradual decay, although both slow and rapidly progressive forms have been reported, exhibiting a large variation in the rate of AD progression (Drachman et al. 1990). While remarkable progress has been gained in the investigation of neurochemical processes accompanying AD, their role in neural degeneration, the main pathological feature of AD, is yet unclear (Selkoe 1987, Kosik 1991). This work is motivated by recent investigations studying in detail the neurodegenerative changes accompanying AD, on a neuroanatomical level. Following the paradigm that cognitive processes can be accounted for on the neural level, we examine the effect of these neurodegenerative changes within the context of a neural network model. This allows us to obtain a schematic understanding of the clinical course of AD.

Neuroanatomical investigations in AD patients demonstrate a considerable decrease in the synapse to neuron ratio, due to *synaptic deletion* (Davies et al. 1987, Bertoni-Freddari et al. 1990). *Synaptic compensation*, manifested by an increase of the synaptic size, was found to take place concomitantly, reflecting a functional compensatory increase of synaptic efficacy at the initial stages of the disease (Bertoni-Freddari et al. 1988, Bertoni-Freddari et al. 1990, DeKosky and Scheff 1990). The combined outcome of these counteracting synaptic degenerative and compensatory processes can be evaluated by measuring the total synaptic area per unit volume (TSA), which was shown to correlate with the cognitive function of AD patients (DeKosky and Scheff 1990).

Our model, presented in section 2, serves as a framework for examining the interplay of synaptic deletion and compensation. This attractor neural network (ANN) is not supposed to represent any specific neuronal tissue, yet we believe that our results are relevant to a large class of neural systems. Deletion is carried out stochastically by removing the fraction d of all synaptic weights. Compensation is modeled by multiplying all remaining synaptic weights by a common factor c . The TSA value is proportional to $c(1 - d)$. Varying c as a function of d specifies a compensation strategy. We assume that the network’s failure rate, measured by the fraction of memories erroneously retrieved, represents the degree of ‘cognitive deficit’ in clinical observations. Reviewing the pertaining pathological and clinical data, we show in section 3 how our model can account for the variability observed in the

clinical course of AD. Our results are further discussed in section 4.

2 The Model

Concentrating on memory degradation, a clinical hallmark of AD (Adams and Victor 1989), we use as our theoretical framework a neural network model of associative memory. Our model is based on the biologically-motivated variant of Hopfield's model (1982), proposed by Tsodyks & Feigel'man (1988). In an ANN, the stored memories are attractors of the network's dynamics, such that when memory retrieval is modeled then, starting from an initial condition sufficiently similar to one of the memory patterns, the network flows to a stable state identical with that memory. The appeal of attractors, as corresponding to our intuitive notion of the persistence of cognitive concepts along some temporal span, has been fortified by numerous studies testifying to the applicability of ANNs as models of the human memory (for a review see Amit (1989)), and also supported by biological findings of delayed, post-stimulus, sustained activity (Fuster 1982, Miyashita and Chang 1988).

All N neurons in the network have a uniform positive threshold T . Each neuron is described by a binary variable $S = \{1, 0\}$ denoting an active (firing) or passive (quiescent) state, respectively. $M = \alpha N$ distributed memory patterns ξ^μ are stored in the network. The elements of each memory pattern are chosen to be 1 (0) with probability p ($1 - p$) respectively, with $p \ll 1$. The weights of the synaptic connections are

$$W_{ij} = \frac{1}{N} \sum_{\mu=1}^M (\xi_i^\mu - p)(\xi_j^\mu - p). \quad (1)$$

The updating rule for neuron i at time t is given by

$$S_i(t) = \Theta \left(\sum_j W_{ij} S_j(t-1) - T \right), \quad (2)$$

where Θ is the step function. The performance of the network is measured by the activities of the memories, as defined by the *overlaps* m^μ ,

$$m^\mu(t) = \frac{1}{p(1-p)N} \sum_{i=1}^N (\xi_i^\mu - p) S_i(t). \quad (3)$$

As shown in the Appendix, there exists an optimal value of the threshold $T = p(1-p)(1-2p)/2$ which ensures the best performance of the network.

Starting with such a memory model we introduce synaptic deletion by randomly deleting some of the incoming synapses of every neuron, leaving each neuron with $l = (1-d)N$

input connections, where $d < 1$ is the *deletion factor*. Synaptic compensation is modeled by multiplying the weights of the remaining synaptic connections by a uniform *compensation factor* $c > 1$. This changes the dynamics of the system to

$$S_i(t) = \Theta \left(\sum_{j \in D_i} cW_{ij}S_j(t-1) - T \right), \quad (4)$$

where D_i denotes a random set of indices corresponding to neurons to which the i -th neuron is connected, and $|D_i|/N = 1 - d \leq 1$. T remains the same value as before.

In the network's 'pre-morbid' state, the memories have maximal stability, achieved by choosing the optimal threshold T that maximizes the increase of the overlap (say m^1), as shown in the Appendix. When the network is initialized with an input pattern which is a corrupted version of one of the stored memory patterns, e. g. ξ^1 , it will flow dynamically into the attractor given by this memory.

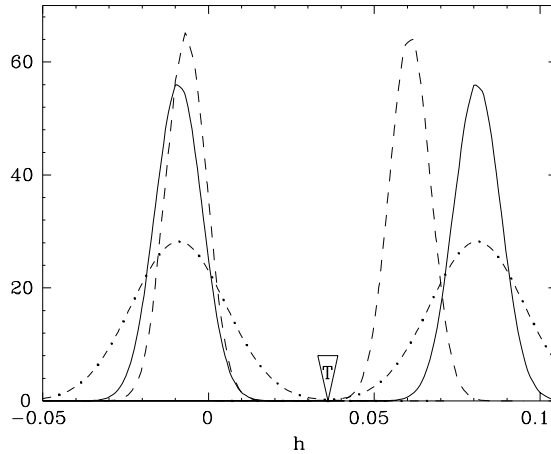


Figure 1: The distribution of the postsynaptic potential ($p = 0.1, \alpha = 0.05$). A. (full line) Initial state: two Gaussian distributions peaked at $-p^2(1-p)$ and $p(1-p)^2$. The *optimal threshold* $T = p(1-p)(\frac{1}{2} - p)$ lies in the middle between the two Gaussian mean values. B. (dashed curve) After deletion ($d = 0.25$), the new peaks of the postsynaptic potential are no longer equidistant from the threshold T . C. (dot-dashed curve) The OPC strategy restores the initial mean values of the postsynaptic potential.

To obtain an intuitive notion of the network's behavior when synaptic deletion and compensation are incorporated consider figure 1. The neurons that stand for firing neurons in the stored memory, and the neurons that stand for quiescent neurons in the stored memory, have distinct post-synaptic potential distributions (the full curves in figure 1). When synaptic deletion takes place, the mean values of the neurons' post-synaptic potential change, and the threshold is no longer optimal (see dashed curves in figure 1). Multiplying

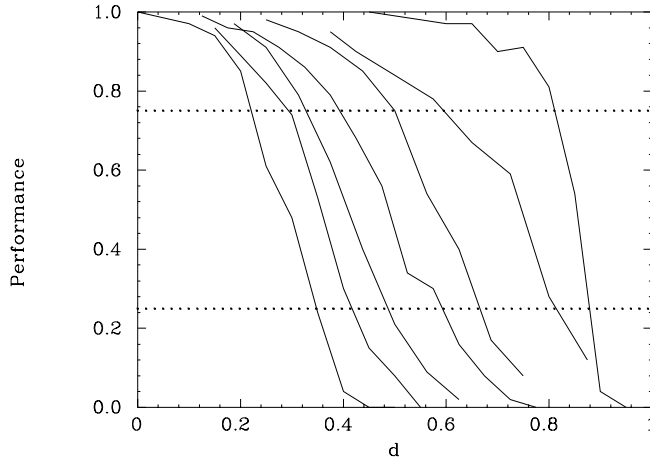


Figure 2: Performance of a network with fixed k compensation. Starting from an initial state that is a corrupted version ($m^1(0) = 0.8$) of a stored memory pattern, we define performance as the percentage of cases in which the network converged to the correct memory. The simulation parameters are $N = 800$ neurons, $\alpha = 0.05$ and $p = 0.1$. The curves represent (from left to right) the performance of fixed strategies with increasing k values, for $k = 0, 0.25, 0.375, 0.5, 0.625, 0.75, 1$. The horizontal dotted lines represent performance levels of 25% and 75%.

the weights of the remaining synaptic connections by an *optimal performance compensation factor* (OPC) $c = \frac{1}{1-d}$, restores the original mean values of post-synaptic potential, and the optimality of the threshold (dot-dashed curves in figure 1). The accompanying increase in the variance of the post-synaptic potential, which is $\sqrt{\frac{1}{1-d}}$ times larger than the original one, leads however to performance deterioration. This is further elucidated in the Appendix.

We can interpolate between the case of deletion without compensation and the OPC within a class of compensatory strategies, defined by

$$\hat{W}_{ij} = cW_{ij}, \quad c = 1 + \left(\frac{1}{1-d} - 1\right)k = 1 + \frac{dk}{1-d}, \quad (5)$$

with the parameter $0 \leq k \leq 1$. All the *fixed* k strategies, examined via simulations measuring the performance of the network at various deletion and compensation levels, display a similar sharp transition from the memory-retrieval phase to a non-retrieval phase, as shown in figure 2. Varying the compensation magnitude k merely shifts the location of the transition region. This sharp transition is similar to that reported previously in cases of deletion without compensation in other models (Canning and Gardner 1988, Koscielny-Bunde 1990).

Figure 3 describes the transition region as a map in the (k, d) plane. The performance levels read off figure 2 delineate the domain over which deterioration occurs. Staying close to the upper boundary of this domain, defines a compensation strategy which enables the

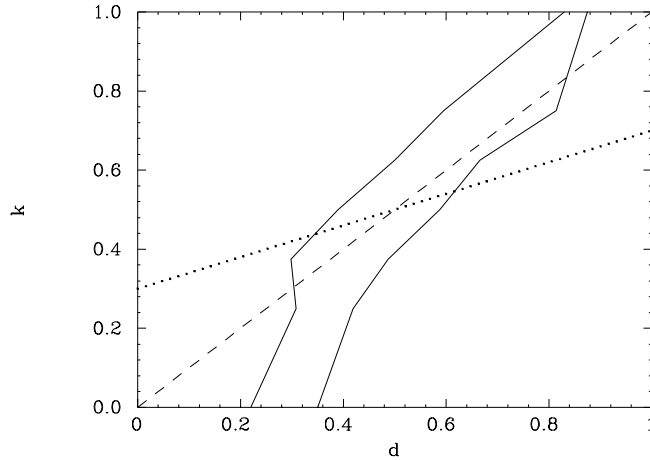


Figure 3: The critical transition range in the (k, d) plane. The full curves represent performance levels of 75% and 25%, derived from figure 2. The straight lines describe the variations employed in two variable compensations presented in figure 4.

system to maintain its performance, with a much smaller amount of synaptic strengthening than that required by the OPC strategy. In face of restricted compensatory resources, such an *optimal resource compensation* strategy (ORC) could be of vital importance. The essence of such ORC strategy is that k is varied as synaptic deletion progresses, in order to retain maximal performance with minimal resource allocation. In figure 4, we present the performance of two *variable k* compensation strategies, which we propose to view as expressions of (albeit unsuccessful) attempts of maintaining an ORC. These examples, indicated on figure 3, include a ‘gradually decreasing’ strategy defined by the variation $k = 0.3 + 0.4d$, and the ‘plateau’ strategy defined by the variation $k = d$. The analogs of these strategies can be found in clinical observations, as shown in the next section where we review the biological and clinical evidence relevant to our model.

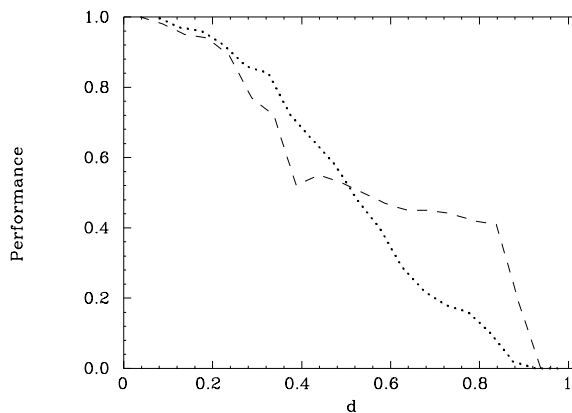


Figure 4: Performance of a network with gradually decreasing (dotted curve) and plateau (dashed curve) compensation strategies.

3 Clinical motivation and implications

As mentioned in the introduction, while synaptic degeneration occurs, the TSA stays constant in some cortical layers at the initial stages of AD. Qualitatively similar synaptic changes have been observed during normal physiological aging, but with significantly lower deletion (Bertoni-Freddari, Meier-Ruge and Ulrich 1988, Bertoni-Freddari et al. 1990). Hence a plausible general scenario seems to involve some initial period of OPC. As AD progresses, synaptic compensation no longer succeeds in maintaining the TSA (Bertoni-Freddari et al. 1990, DeKosky and Scheff 1990). In advanced AD cases, severe compensatory dysfunction has been observed (Buell and Coleman 1979, Flood and Coleman 1986, DeKosky 1990).

Young AD patients are likely to have high compensation capacities, and therefore can maintain an OPC strategy ($k = 1$, in figure 2) throughout the course of their disease. This will then lead to a rapid deterioration when the reserve of synaptic connections has been depleted. Indeed, young AD patients have been reported to have a rapid clinical progression (Heston et al. 1981, Heyman et al. 1983), accompanied by severe neuronal and synaptic loss (Hansen et al. 1988). A similar clinical pattern of rapid memory decline, already manifested with less severe neuroanatomical pathology, was found in very old patients (Huff et al. 1987). We propose that in these old patients, the rapid clinical decline results from the lack of compensatory capacities ($k = 0$, in figure 2), possibly of the kind observed by Buell and Coleman (1979) and Flood and Coleman (1986).

Rapid cognitive decline characterizes a minority of AD patients. Most patients show a continuous gradual pattern of cognitive decline (Adams and Victor 1989, Katzman 1986, Katzman et al. 1988), taking place along a *broad span* of synaptic deletion (DeKosky and Scheff 1990). As shown in figure 2, this performance decline cannot be accounted for by any network employing fixed k compensation. Variable compensation, such as that defined by the gradually decreasing strategy, is needed to explain the memory decline observed in the majority of AD patients, as shown in figure 4. The clinical state of some AD patients remains stable at mild to moderate levels for several years before finally rapidly decreasing (Cummings and Benson 1983, Katzman 1985, Botwinick, Storandt and Berg 1986). This can be accounted for by a ‘plateau’ strategy whose performance, shown in figure 4, stays at an approximately constant level over a large domain of deletion.

Synaptic deletion and compensatory mechanisms play a major role also in the patho-

genesis of Parkinson disease (Zigmond et al. 1990, Zigmond and Calne 1991). The significant incidence of AD patients having accompanying extra-pyramidal parkinsonian signs (Mayeux, Stern and Spanton 1985, Stern et al. 1990), naturally raises the possibility that such patients may have a decreased synaptic compensatory potential in general (Horn and Ruppin 1992). The cognitive deterioration of these AD patients is faster than that of AD patients without extra-pyramidal signs. This fits well with our proposal that severely deteriorated synaptic compensation capacity leads to an accelerated rate of cognitive decline in AD patients. This issue is still inconclusive because the PD-AD combination may be a specific syndrome on its own.

4 Discussion

In accordance with the findings that neuronal loss in AD is less than 10% even at advanced stages (Katzman 1986), and that the synapse to neuron ratio is significantly decreased (Davies et al 1987, Bertoni-Freddari et al. 1990), we have concentrated on studying the role of the synaptic changes. Simulations we have performed incorporating neuronal loss have shown similar results to those presented above. We conclude therefore that the important factors are indeed the number of synapses retained and the compensation strategy employed, whose interplay may lead to various patterns of performance decline.

As any current neural model of human cognitive phenomena, our model necessarily involves many simplifications. The TF formal neurons are obviously a very gross simplification of biological neurons. As in most Hopfield-like ANNs, the network has no spatially specified architecture. For clarity of exposition of our main ideas, we have assumed that all compensation strategies are applied uniformly to all retained synapses. Our analysis also holds for non-uniform compensation, i.e., when each remaining synaptic weight is multiplied by a random variable with mean value c and variance σ^2 , since the same averages of the postsynaptic potentials are obtained (see Fig. 1 and the Appendix). Obviously, if the variance is too large, then no compensation strategy can be conceived of any more. Motivated by the biological evidence testifying to the sparsity of neural firing (Abeles *et al.* 1990), we have assumed a relatively small fraction p of firing neurons. Simulations performed with higher p values (e.g. 0.2) indicate that the results remain qualitatively the same. However, it should be noted that as p is increased the approximation of the network's overlap dynamics presented in the Appendix becomes less and less accurate.

The variable compensation strategies which we have discussed rely on the fact that there is some span in the (k, d) plane over which deterioration takes place, as shown in figure 3. As N is increased, the width of the domain over which deterioration occurs keeps getting narrower, thus limiting the possibilities of maneuvering between deletion and compensation. Hence, one may claim that our conclusions, which are based on simulations of small scale networks, do not hold for the brain.

One possible answer to this problem is that there may exist important modules in the brain whose function depends on the correct performance of just some thousands of neurons (Eccles 1981). For large cortical modules, this objection may be resolved by considering the effect of noise present in the brain. To account for the latter, any realistic paradigm of memory recall should entail the recognition of a spectrum of noisy inputs presented to the network. Figure 5 displays the performance of the network in the (k, d) plane obtained via simulations with two distinct initial overlap values ($m^1(0) = 0.8$ and $m^1(0) = 0.95$), together with the theoretical results for the infinite N limit. These results show that even in this limit, the corresponding performance curve always retains a finite width as long as the network processes input patterns with a broad range of initial overlaps. Consequently, the realization of variable compensatory strategies may indeed be feasible in the brain.

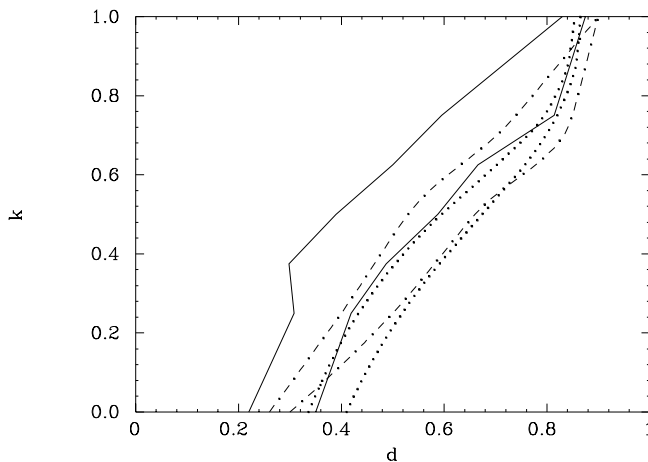


Figure 5: The critical transition range in the (k, d) plane. The full curves represent performance levels of 75% and 25%, with initial overlap $m(0) = 0.8$ (identical to figure 2). The dash-dotted curves represent performance levels of 75% and 25%, for initial overlap $m(0) = 0.95$. The dotted curves represent the theoretical results which follow from the analysis presented in the Appendix, delineating the estimates of when the corresponding basins of attraction cease to exist in the infinite N limit. These curves lie close to the 25% lines of the simulations.

The decline in the network's performance resulting from synaptic deletion is coupled with a decrease in the network's overall activity. This observation gives rise to the possibility that, although being defined 'globally' as 'strategies', synaptic compensation may take place via local feedback mechanisms. Accordingly, the decreased firing rate of a neuron being gradually deprived of its synaptic inputs may enhance the activity of cellular processes strengthening its remaining synapses. This scenario seems to lead to fixed OPC compensation in a rather straightforward manner, but as synaptic deletion may be non-homogeneous the effects of the resulting spatially non-uniform compensatory changes should be further investigated. The non-vanishing width of the (k, d) -plane transition range shown above is essential for the feasibility of an ORC strategy, so that local mechanisms can 'trace' the decreasing performance and 'counteract' it before the performance collapses entirely.

Finally let us comment on possible examinations and applications of our model. An ideal experiment would involve a series of consecutive measurements of synaptic strength and cognitive abilities. In the light of obvious difficulties concerning such tests, we may have to resort to comparing biopsies and autopsies, as in DeKosky and Scheff (1990), preferably on the same patients. Our model demonstrates the importance of maintaining the TSA for the preservation of memory capacity and, therefore, mental ability of AD patients. This may suggest that future therapeutic efforts in AD should include an attempt to mobilize compensatory mechanisms facilitating the regeneration of synaptic connectivity.

Appendix

Qualitative features of our model can be derived from a simple analysis of the first iteration step. Starting with a state which is close to ξ^1 with overlap $m(0)$ we wish to find whether the network flows into the correct memory. Using the dynamics defined in the text we find for $t = 1$

$$S_i(1) = \Theta \left[\frac{1}{N} c \left(\sum_{j \in D_i} (\xi_i^1 - p)(\xi_j^1 - p) S_j + \sum_{\mu > 1} \sum_{j \in D_i} (\xi_i^\mu - p)(\xi_j^\mu - p) S_j \right) - T \right] \\ = \Theta [c(1-d)(\xi_i^1 - p)p(1-p)m(0) + \mathbf{N} - T] \quad (6)$$

where we have separated the signal from the noise term. The latter has zero average, $\langle \mathbf{N} \rangle = 0$, and variance of $\langle \mathbf{N}^2 \rangle = c^2(1-d)p^2(1-p)^2\alpha s(0)$, where $s(0) = P(S_i(0) = 1) = 1 - p - m(0) + 2pm(0)$. In view of the Gaussian noise term we write the probability in terms of an error-function,

$$P(S_i = 1) = \text{erf} \left(\frac{c(1-d)(\xi_i^1 - p)p(1-p)m(0) - T}{\sqrt{c^2(1-d)\alpha s(0)p^2(1-p)^2}} \right). \quad (7)$$

This results in the following expression for the first iteration:

$$m^1(1) = \frac{1}{p(1-p)} [(1-p)P(\xi_i^1 = 1)P(S_i(1) = 1|\xi_i^1 = 1) - pP(\xi_i^1 = 0)P(S_i(1) = 1|\xi_i^1 = 0)] = \quad (8) \\ = P(S_i(1) = 1|\xi_i^1 = 1) - P(S_i(1) = 1|\xi_i^1 = 0) = \\ \text{erf} \left(\frac{(1-p)m^1(0)p(1-p)c(1-d) - T}{\sqrt{\alpha c^2(1-d)s(0)p^2(1-p)^2}} \right) - \text{erf} \left(\frac{(-p)m^1(0)p(1-p)c(1-d) - T}{\sqrt{\alpha c^2(1-d)s(0)p^2(1-p)^2}} \right).$$

In the limit $m(0) \rightarrow 1$ one finds the maximal value of equation 8 to be obtained for the following choice of the optimal threshold

$$T^* = c(1-d)p(1-p)(1-2p)/2. \quad (9)$$

For $c = 1$ and $d = 0$ this coincides with the choice we have made. Moreover, as long as

$$c = \frac{1}{1-d}, \quad (10)$$

which was defined as the OPC strategy, T remains optimal. This fact was expressed graphically in figure 1. The two Gaussian distributions in this figure correspond to the two terms in equation (8).

In the simulations we have looked for the cases in which the system converged onto the correct fixed points. This involves iterating the equations of motion, which is in general different from iterating expressions like equation (8) because of possible correlations between the different time steps. Nonetheless we may think of the iteration of equation (8) (replacing the labeling 0 and 1 by $n-1$ and n) as a rough estimate for the strongly diluted infinite system (Evans 1989). Starting out with different values for $m(0)$ we find the dotted curves in figure 5 which show the border-lines between convergence and non-convergence to the correct memory.

An alternative to this derivation is to use the replica symmetry assumption. We have carried out such an analysis (Herrmann et al. 1992). The results are similar, though not identical, to the ones reported above.

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