How training and testing histories affect generalization: a test of simple neural networks

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We show that a simple network model of associative learning can reproduce three findings that arise from particular training and testing procedures in generalization experiments: the effect of (i) ‘errorless learning’, (ii) extinction testing on peak shift, and (iii) the central tendency effect. These findings provide a true test of the network model which was developed to account for other phenomena, and highlight the potential of neural networks to study the phenomena that depend on sequences of experiences with many stimuli. Our results suggest that at least some such phenomena, e.g. stimulus range effects, may derive from basic mechanisms of associative memory rather than from more complex memory processes.

Keywords: animal behaviour; neural networks; range effects; generalization

1. INTRODUCTION

This paper deals with a general issue in the study of animal behaviour that we call path dependence. The expression refers to the fact that different histories of experiences (paths) may at first seem to produce the same behavioural effects, yet reveal important differences when further examined. For instance, two training procedures may establish the same discrimination between the two training stimuli, yet produce different responses to other stimuli, because the two paths have produced different internal states within the animal. There are several reasons why path dependence is an important issue. First, it comprises many phenomena that can provide stringent tests for theories of behaviour. Second, path dependence is at the root of several controversies; for instance, whether animals encode absolute or relative characteristics of stimuli (Spence 1936; Helson 1964; Thomas 1993; Sarris 2003), whether learning phenomena such as backward blocking and unovershadowing imply, in addition to basic associative learning, stimulus–stimulus associations or changes in stimulus associability (Wasserman & Berglan 1998; Le Pelley & McLaren 2003; Ghirlanda 2005).

In this paper, we use a simple neural network model of basic associative learning (Blough 1975; Enquist & Ghirlanda 2005) to show how path dependence can arise from fundamental properties of associative memory. The model has two core components: (i) distributed representations of stimuli based on the knowledge of sensory processes and (ii) a simple learning mechanism that can associate stimulus representations with the responses. We consider examples of path dependence in experiments on generalization (or ‘stimulus control’). These consist of a training phase in which animals are trained to perform a specific response to several stimuli and a test phase in which responding to a set of stimuli is recorded. The test stimuli often lie on a ‘stimulus dimension’ such as light wavelength or object size so that generalization is often described as a response gradient over the dimension. In generalization experiments, path dependence appears as differences in the shape of generalization gradients; different paths correspond to different training or testing procedures. We show that the model accounts for the following phenomena (see §3 for details): lack of peak shift after ‘errorless discrimination learning’, decrease of peak shift during extinction testing and the shift of generalization gradients towards the average of the test stimuli (a kind of range effect).

Although we consider laboratory experiments, it is hardly necessary to note that path dependence exists in the wild as well, where paths are the consequence of environmental events rather than being arranged by an experimentalist.

2. MODEL

(a) The neural network

Nervous systems can be seen as a flexible structure that can be programmed to generate almost any behaviour, e.g., relationships between stimuli and responses. Concretely, such programming includes both the formation of the neural network and its pattern of connectivity and the adjustment of connections between cells in the network. Neural network models provide an understanding of how such processes can ultimately produce the behaviour we see in animals (Arbib 2003; Enquist & Ghirlanda 2005). Previous work has shown that the simple neural network...
models of associative learning can reproduce many fundamental findings of learning and generalization (Blough 1975; Enquist & Ghirlanda 2005; Ghirlanda 2005). Here, we use a standard feed-forward network with an array of input nodes connected directly to one output node (there are no hidden nodes). Stimuli are modelled as eliciting graded patterns of activity in the array of input nodes. We write $S_i$, the activity induced in input node $i$ by stimulus $S$ ($i=1, \ldots, N$). The input nodes are connected to the output node by weighted connections, the weight attached to node $i$ being $W_i$. The strength or likelihood of responding to $S$ is assumed to be an increasing function of the weighted sum $r_S$

$$r_S = \sum_i W_i S_i.$$  

(2.1)

(b) Learning

We model learning by the so-called $\delta$ rule (McClelland & Rumelhart 1985), a simple case of gradient descent algorithms (Haykin 1999) that was first derived by Widrow & Hoff (1960) and introduced to animal learning theory by Blough (1975), based on previous work by Rescorla & Wagner (1972). At each stimulus presentation, the algorithm prescribes a change $\Delta W_i$ in weight $W_i$ according to

$$\Delta W_i = \alpha(\lambda - r_S) S_i,$$  

(2.2)

where $\lambda$ is the maximum value responding to $S$ can attain given the applied reinforcer, and $\alpha$ mainly regulates the speed of learning (Widrow & Stearns 1985). Equation (2.2) is capable, through repeated applications, of establishing a different response to each of many stimuli, provided the corresponding patterns of activity satisfy certain requirements (‘linear separability’). We refer to the literature for technical details and other applications to behaviour theory (Blough 1975; Widrow & Stearns 1985; McClelland & Rumelhart 1986; Rumelhart & McClelland 1986; Haykin 1999; Enquist & Ghirlanda 2005). Weights are assumed to start from a value of zero (drawing weights at random from a distribution symmetrical around zero would lead to the same conclusions).

(c) Model stimuli and generalization tests

The results below do not depend on the precise details of how stimuli are modelled, as long as the following general properties hold: (i) input node activity is positive; (ii) each stimulus corresponds to a graded pattern of activity in input nodes; (iii) physically more similar stimuli correspond to more similar patterns of activity; and (iv) higher intensity of stimulation corresponds to higher input node activity. Figure 1 shows the kind of model stimuli we use in practice. Dimensions such as sound frequency, light wavelength or spatial position can be modelled by translating an activity profile like the one in figure 1b over the input array of the network. This simple scheme captures the empirical observation that a stimulus change along these dimensions causes a change in the pattern of receptor activity but not in total activity. Conversely, stimulus intensity dimensions are modelled by increasing or decreasing input node activity without changing.
called \( S^+ \), and to ignore another one, called \( S^- \). This may be achieved by instrumental conditioning, whereby responses to \( S^+ \) are ‘reinforced’, e.g. with food, whereas responses to \( S^- \) are not (Pearce 1997). Details of training may vary, but it is most common to first train the desired response to \( S^+ \) and then to introduce \( S^- \). The animal will usually respond to \( S^- \) in the first stages of discrimination training (especially if \( S^- \) is similar to \( S^+ \)), but if such responses are never reinforced, the animal will respond less and less to \( S^- \). In practice, discrimination training is continued until a criterion is met such as ‘three times more responding to \( S^+ \) than to \( S^- \) or ‘no responses to \( S^- \) in a 10 min period’.

When a generalization test is performed after discrimination learning, one may find that the stimulus which elicits most responses is not \( S^+ \), but a stimulus that is displaced away from \( S^+ \) so as to be more different from \( S^- \) (figure 2a, dotted line). Since its discovery by Hanson (1959), this phenomenon, called the ‘peak shift’, has fuelled extensive research to understand how a stimulus that was never reinforced (and in many cases, never experienced) could be more powerful in eliciting a response than a reinforced stimulus (Mackintosh 1974). It has been known, at least since the pioneering work of Blough (1975), that simple network models can reproduce the peak shift phenomenon. We will briefly review the mechanism below, referring to Enquist & Ghirlanda (2005) for further discussion. The dotted line in figure 2b shows a peak shift obtained from a neural network simulation of a discrimination experiment. For technical details regarding this and all other simulations in the paper, we refer to the electronic supplementary material.

Figure 2a also shows a generalization gradient without a peak shift (continuous line), although the same \( S^+ \) and \( S^- \) have been used in training. The difference is that training did not follow the standard procedure outlined above but an alternative one, ‘errorless discrimination learning’, developed by Herbert Terrace at the beginning of the 1960s. In errorless discrimination learning, \( S^- \) is introduced gradually rather than abruptly. For instance, in the experiment in figure 2, Terrace (1964) trained a discrimination between a 580 nm monochromatic light (\( S^+ \)) and a 540 nm light (\( S^- \)), beginning with a very faint 540 nm light whose intensity was progressively increased until an intensity equal to \( S^+ \) was reached. The name ‘errorless learning’ derives from the fact that the animal responds very little to \( S^- \) throughout training. Intuitively this happens because, at any given moment, \( S^- \) is very similar to previously unreinforced stimuli (including, at the start of training, the experimental background), and thus has only a small probability of eliciting a response. Additionally, initial \( S^- \) presentations are so brief that the animal is effectively prevented from responding.

Terrace’s finding that errorless learning prevents the peak shift has been replicated a few times, but no agreement exists as to its causes (Purtle 1973). Our aim is to explore what insight can be gained by simulating errorless learning with neural networks. We mimic Terrace’s procedure by starting with a model \( S^- \) of low intensity (low activation of network input nodes) and progressively increasing its intensity, as shown for instance in figure 1d. The resulting generalization gradient (continuous line in figure 2b) peaks on \( S^+ \) in agreement with Terrace’s empirical result. To understand why this happens, we plot in figure 3 the weight values obtained after both standard and errorless training. After standard training (dotted line), both positive and negative weights develop, associated respectively with parts of the input array most activated by \( S^+ \) and \( S^- \). When peak shift occurs, maximum response is observed for stimuli that are close to \( S^+ \), but more distant from \( S^- \) than \( S^+ \) itself. Such stimuli retain most of \( S^+ \)’s ability to excite nodes with positive weight while activating nodes with negative weights significantly less, which results in a more favourable balance between excitation and inhibition. During errorless learning, on the other hand, the input nodes most stimulated by \( S^- \) develop very small negative weights (continuous line). Thus, the gains of departing from \( S^- \) cannot offset the losses caused by departing from \( S^+ \). Interestingly, this explanation is consistent with Terrace’s suggestion that errorless learning results in little inhibition being associated with \( S^- \).

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The reason why the weights develop as shown in figure 3 can be understood by imagining what happens in the initial phases of training. The low intensity $S^-$ used at the start of errorless learning is very similar to the background stimuli to which response is low (e.g. the dark response key in Terrace’s experiment). To ensure that response to such an $S^-$ be low, therefore, it is sufficient to adjust the weights only a little. On the other hand, at the beginning of standard training, $S^-$ is an intense stimulus not unlike $S^+$ and thus produces ‘errors’ in the form of high network output, while the desired response is a low output. The learning algorithm must thus decrease response to $S^-$ considerably, which is achieved by attracting negative weight to the input nodes most stimulated by $S^-$.  

(b) Disappearance of peak shift in extinction

The outcome of a generalization test can be affected by different aspects of the testing procedure, e.g. its duration and what stimuli are used. The reason is, of course, that animals continue to learn during a test. Thus test results are not, as one would like, simply the result of probing the animal, but are partly due to learning caused by probing itself. The most common testing paradigm is testing ‘in extinction’, i.e. by unreinforced presentation of test stimuli. This causes a generalized decrease in response and can also change the shape of the generalization gradient. An interesting finding that we consider here is the reduction of peak shift during testing in extinction (figure 4a; see also Purtle 1973).

To model this finding, we teach the network a discrimination between $S^+$ and $S^-$, then run a first generalization test, which shows a peak shift (figure 4b, dotted line). We then continue to test mimicking the extinction procedure, i.e. we apply the δ rule after each stimulus presentation with a low target value (low $\lambda$ in equation (2.2)). In the generalization gradient produced after many such presentations, we find a greatly reduced peak shift. Network weights at the beginning and end of extinction testing are shown in figure 5, where it is apparent that extinction testing has reduced the difference between the positive and the negative weights which underlies peak shift. Testing in extinction has also reduced the absolute values of the weights, which results in a general decrease in network output that parallels the decrease in response as observed in the experiments.

(c) Range and frequency effects

Any set of stimuli may be, in principle, used in a generalization test. It is most common to use a range of evenly spaced stimuli roughly centred around the training ones, with the aim of getting an unbiased picture of generalization (with only partial success, as seen above). However, different kinds of tests have been used specifically to study how generalization is affected by post-training experiences. The most common manipulations include presenting some test stimuli more often than others and using only stimuli within a restricted range (for example, reviewed in Thomas et al. 1992). The changes in generalization gradients brought about by such procedures may be collectively labelled as ‘range and frequency effects’ and have generated a considerable debate about the underlying memory processes (Spence 1936; Helson 1964; Parducci 1965; Thomas 1993; Sarris 2003).

One common finding is that extensive testing causes a ‘central tendency effect’, whereby the generalization gradient appears to be shifted towards the middle of the stimulus range used in the test (figure 6a; reviewed by Thomas et al. 1992). It is easy to test the network in these conditions, simply running the tests in extinction with different ranges of stimuli. The outcome of such tests is indeed a central tendency effect (figure 6b). Figure 7 shows how testing with different ranges of stimuli modifies the weight array: probing with a particular stimulus range causes a shift in the weight array towards the middle of the range.

Our results suggest that at least some range and frequency effects may arise from simple mechanisms of associative learning, while current thinking often appeals to more complex memory processes (cf. ‘adaptation level’ theory and ‘frequency-range’ theory;
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Figure 6. Central tendency effect. (a) Generalization gradients obtained in extinction with different ranges of test stimuli, indicated by the lines below the graph in matching style, after identical training to respond to $S^*$. The gradient peak appears displaced towards the centre of the test range (data from Thomas & Jones 1962, asking humans to identify a 525 nm light). (b) Network simulation of the same experiment, showing a similar central tendency effect.

Figure 7. Weight arrays for the simulations in figure 6a. The grey line represents weights just after training, the other lines show how weights have changed after testing in extinction with three stimulus ranges. In addition to a general decline caused by the extinction procedure, we see a shift in the pattern of weights such that the largest weights move toward the centre of the probed stimulus range.

Helson 1964; Parducci 1965; Thomas et al. 1992; Sarris 2003. The need for such additional processes is partly inferred from the belief that range and frequency effects are virtually absent in non-human animals, although it is possible to find examples in the animal literature (reviewed by Thomas 1993; Sarris 2003). Perhaps, the relative ease with which range and frequency effects appear in humans may just follow from the fact that humans learn faster (i.e. the effect of testing is seen even in relatively short tests).

4. DISCUSSION
In this exploratory study, we have used simple neural networks to test the hypothesis that path dependence phenomena arise from basic mechanisms of associative learning in distributed memory systems. We have shown that a simple network model of learning can reproduce three particular findings: the effect of errorless learning and of extinction testing on peak shift and the central tendency effect. We chose to consider these findings for several reasons. A first one is that they provide a true test of neural network models, which were developed to account for different phenomena. We stress that we have used, without modification, a very basic model, essentially Blough’s (1975) model with the addition that stimulus representations be built with knowledge of relevant sensory processes (Ghirlanda & Enquist 1999; Enquist & Ghirlanda 2005).

A second reason is that the considered phenomena have been known for many decades, yet their theory is still unsatisfactory. The effects of errorless learning and extinction testing have been repeatedly considered in the peak shift literature (Purtle 1973; Mackintosh 1974), but the theory is limited to verbal arguments such that the animals may learn ‘from the experience of being tested’ (Prokasy & Hall 1963, quoted by Purtle 1973). Range and frequency effects have received considerable attention (Thomas et al. 1992; Thomas 1993; Sarris 2003), but the extent to which they can be accounted in terms of simple associative learning is still unknown. The main theoretical difficulty posed by all these phenomena, and by path dependence in general, is that they require us to track the cumulative effect of sequences of experiences with many stimuli. This is difficult in most models (to put it mildly) and reveals one crucial advantage of neural networks: the ability to simulate arbitrary sequences of experiences and to get predictions about responding to any stimulus that can be received. Simple neural networks are also amenable, in some cases, to mathematical analysis, although we have not pursued this approach here (see Haykin 1999; Enquist & Ghirlanda 2005).

In conclusion, neural networks provide a very natural framework to study path dependence and thus increase our knowledge of how experiences shape behaviour. The neural network models are already a promising account for a large body of behavioural phenomena (Enquist & Ghirlanda 2005) and including path dependence would contribute to a unified picture of how nervous systems bring about behaviour.

REFERENCES


