Recalling an aversive experience by day-old chicks is not dependent on somatic protein synthesis

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The translation of a short-term experience into longer term memory (consolidation) requires protein synthesis, presumed to be necessary for the rescultping of synapses (Hebb 1949). Inhibitors of protein synthesis, administered around the time of training or 4–6 h later, produce lasting amnesia for the task (Davis and Squire 1984; Rose 2000). Beyond this time, the memory is insensitive to the inhibitors and has been regarded as permanent (long-term memory). However, recently reconfirmed older observations show that reminding the animal of the previously learned experience renders the memory labile once more (Sara 2000a,b; Nader 2003; Dudai 2004). Administration of protein synthesis inhibitors in association with the reminder for an aversive experience produces amnesia for the task, in some cases apparently permanent (Nader et al. 2000; Nader 2003), in others more transient (Litvin and Anokhin 2000; Milekic and Alberini 2002; Eisenberg and Dudai 2004). This has prompted an ongoing debate, i.e., is the amnesia due to a blockade of the same biochemical cascade as is involved in the initial consolidation (hence, permanently preventing “reconsolidation”), or does it represent a temporary failure to access the memory (retrieval) (Nadel and Land 2000; Alberini 2005). Of course, in some senses this distinction is artificial, as any reminder inevitably constitutes a new experience and will involve some learning, which may be part of a process leading to extinction of the earlier memory (Vianna et al. 2001). A further complexity is added by the fact that even without reminder, putative memory traces are not entirely stable, migrating from one brain region to others over a period that may vary from hours to weeks (Myers and Squire 1984; Myers and Morgan 1981). These authors reported that bilateral injections of 5µg (15 nmol) Colchicine into the forebrain shortly after training resulted in a transient amnesia for the avoidance response. We began by confirming that Colchicine injected bilaterally into the IMMP at doses of up to 20 µg was without overt behavioral or toxic effects. Injections of Colchicine at 15 nmol at 30–10 min pre-training were without effect on acquisition of the task or recall in chicks tested at varying times up to 24 h subsequently (data not shown). However, in accordance with Bell and Morgan’s observations, we found that bilateral injections of 15 nmol Colchicine/hemisphere into the IMMP, 15 min post-training, results in amnesia in chicks tested 3 h subsequently. The amnesia is, however, transient; chicks trained, injected with initial training. Furthermore, whereas the biochemical locus of change following training is in the intermediate medial mesopallium (IMMP, previously called IMHV) (Reiner et al. 2004), following a reminder it is in the region we had earlier (Rose 2000) identified as a putative “storage site” for the memory trace, the medial striatum (MS, previously called LPO).

One explanation for the differences in the amnestic effect of Ani could be that while the initial learning experience involves enhanced gene expression and somatic protein synthesis followed by the transport of the newly synthesized proteins to the synapse, re-evoking the experience by way of a reminder engages only local (dendritic/synaptic) protein synthesis. That such synthesis can occur in dendritic spines and presynaptic elements (synaptoneurosomes) is well established (Steward and Worley 2002; Tang and Schuman 2002). We reasoned that if this were the case, then while transiently blocking axonal and dendritic flow during consolidation should result in amnesia for the task, this would not be the case following recall of the experience. Such a transient blockade, lasting minutes to hours, occurs if microtubular structure is disrupted, which can be achieved by administration of Colchicine (Borisy and Taylor 1967a; Edson et al. 1993). In the experiments reported here we have examined the effects of Colchicine on recall for the passive avoidance task following both training and reminder.

Results

Colchicine effect on recall following training

We began by replicating and extending an earlier study by Bell and Morgan (1981). These authors reported that bilateral injections of 5µg (15 nmol) Colchicine into the forebrain shortly after training resulted in a transient amnesia for the avoidance response. We began by confirming that Colchicine injected bilaterally into the IMMP at doses of up to 20 µg was without overt behavioral or toxic effects. Injections of Colchicine at 15 nmol at 30–10 min pre-training were without effect on acquisition of the task or recall in chicks tested at varying times up to 24 h subsequently (data not shown). However, in accordance with Bell and Morgan’s observations, we found that bilateral injections of 15 nmol Colchicine/hemisphere into the IMMP, 15 min post-training, results in amnesia in chicks tested 3 h subsequently. The amnesia is, however, transient; chicks trained, injected with...
further analysis was made using a post hoc least-significant difference (ANOVA), and where significant differences were found between groups, statistical analysis of behavioral data was by analysis of variance (ANOVA), and where significant differences were found between groups, further analysis was made using a post hoc least-significant difference test.

Colchicine as before, and tested 24 h later showed good recall (Fig. 1). If the Colchicine injections were delayed to 30 min post-training, however, no amnesia was apparent (data not shown).

Electron micrographs from brain perfused 30 min after injection of Colchicine into the IMMP showed considerable disruption of microtubules in the region compared with controls (Fig. 2A, B). By 3 h post-injection, the microtubules in Colchicine-injected animals had begun to reform (Fig. 2C). To confirm that Colchicine does indeed also disrupt transport of newly synthesized proteins to the synapses, we injected 14-C-leucine bilaterally into the IMMP, followed 15 min later by 15 nmol Colchicine. Three hours later, the chicks were killed, forebrain regions containing IMMP dissected, and crude synaptoneurosome fractions prepared by centrifugation. The purity of the synaptoneuroosomal preparation was routinely checked by electron microscopy (Fig. 2D). Colchicine did not affect the specific radioactivity of incorporated leucine in the trichloroacetic acid-precipitated proteins in the homogenate, indicating that it was without effect on protein synthesis per se, but reduced the proportion of the de novo synthesized proteins recovered in synaptoneuroosomes (Fig. 2E). Thus, Colchicine does not disrupt protein synthesis, but merely blocks the delivery of the proteins to the synapse. This blockage is transient, as by 3 h, the microtubules in this region are partially reconstituted.

Colchicine effect on recall following a reminder

As Colchicine produces transient amnesia following training, we then asked whether it also does so following a reminder. In our earlier experiments (Anokhin et al. 2002; Salinska et al. 2004) we found that the post-reminder time window at which Ani causes transient amnesia in the chick does not coincide exactly with that post-training; therefore, we divided chicks into two groups. The first was trained, as in Figure 1, injected with Colchicine 15 min later, and tested 1, 2, 3, or 6 h subsequently. Chicks in the second group were trained and remained in their pens for 24 h, after which they were given a reminder by presenting them with a dry chrome bead identical to the one that they had previously pecked and found to taste bitter. Birds that pecked the bead at the reminder (<10%) were excluded at this point. Fifteen minutes later, birds that avoided the bead during the reminder were injected with Colchicine and tested 1, 2, 3, or 6 h subsequently. As Figure 3A shows, birds injected with Colchicine following training showed a transient amnesia, birds injected post-reminder showed normal retention (Fig. 3B).

Anisomycin effect following a reminder

To confirm that, unlike the situation post-training, Ani and Colchicine differed in their effects post-reminder, we compared them directly in the same experiment. Chicks were trained and divided into two groups, and each group then subdivided into three. The first group was injected with saline or Ani (5 min post-training, the time-point used in our earlier experiments) (Salinska et al. 2004) or Colchicine (15 min post-training) and tested 3 h later as in Figure 1. The second group was given a reminder 24 h post-training and injected with Colchicine, Ani, or saline as before. Three hours later, all birds were tested as before. As Figure 4 shows, Ani-injected birds were amnesic following either training (Fig. 4A) or reminder (Fig. 4B), as predicted from our earlier
Colchicine, consolidation, and reconsolidation

Figure 3. Effect of Colchicine on retention following training or reminder. Animals were trained as described in the Materials and Methods and reminded 24 h after training, as described in the text. (A) Chicks were either injected 15 min post-training and tested at 1, 2, 3, or 6 h subsequently, or (B) remained in their pens for 24 h, followed by a reminder and Colchicine injection and tested at 1, 2, 3, or 6 h post-reminder. Results are presented as mean ± SEM. n = 45–60 for each group. (*) \( P < 0.005 \); (**) \( P = 0.0007 \); (*** \( P = 0.000025 \). Statistical analysis of behavioral data was by analysis of variance (ANOVA) and where significant differences were found between groups, further analysis was made using a post hoc least-significant difference test.

Discussion

Colchicine has a variety of effects on intracellular structures (Borisy and Taylor 1967b; Wilson et al. 1999) and by virtue of disrupting microtubules, blocks axonal flow and, hence, the transport of de novo synthesized proteins to the synapse (Gardiol et al. 1999; Steward and Worley 2002). This blockade will include the proteins required for the synaptic restructuring necessary for memory consolidation, such as the cell-adhesion molecules (Mcleunnic et al. 1995; Scholey et al. 1995), which are synthesized in the soma. Blocking axonal flow does not affect the synthesis of these proteins, but results in their accumulation in the soma until such time as the blockade is lifted. The data of Figure 2 confirms this for the chick. In contrast to the well-established finding that, as in other species and tasks, injection of Ani around the time of training in the passive avoidance task in the chick results in lasting amnesia (Davis and Squire 1984; Rose 2000), Colchicine produces transient amnesia. As shown in Figure 1, and as reported previously by Bell and Morgan (1981), following a Colchicine injection, memory, absent at 3 h, is restored by 24 h. Although this temporary blockade might have been a consequence of other effects of Colchicine on cell structure, such as tenascin filaments (Domnina et al. 1995), the most probable interpretation is that only when the microtubular block is lifted are the newly synthesized cell-adhesion molecules transported and, accordingly, enable memory consolidation to proceed. Implicit in this argument is that any putative synaptic "tags" (Frey and Morris 1998) must remain in place during the blockade of transport.

However, in contrast to the effect of Ani, disrupting axonal transport by injecting Colchicine into the IMP does not affect retention following a reminder 24 h later (Fig. 3). One possible explanation for this would be that at 24 h, memory for the task is no longer dependent on the IMP but on the MS, to which there is a projection via the archistriatum (Csillag 1999). Although pre-training lesions of the IMP result in amnesia for the avoidance response, lesions made some 3 h post-training are without amnestic effect (Gilbert et al. 1991). In contrast, pre-training lesions of the MS are not amnestic, while post-training lesions are (see also, Salinska et al. 2004). However, post-reminder injections of Colchicine into the MS, like those into the IMP, were without effect on subsequent recall. Further, Ani continued to produce amnesia even in the presence of Colchicine (Fig. 4), suggesting that in this case it is blocking the synthesis of synaptoneurosomal proteins on mRNA already present. There is good evidence both for the presence of such mRNA and for the functional significance of the proteins whose synthesis it makes possible (Job and Eberwine 2001; Tang and Schuman 2002) as well as for spine plasticity (Carlisle and Kennedy 2005). The failure of Colchicine to produce amnesia following a reminder is thus most straightforwardly explained on the assumption that, in contrast to the somatic protein synthesis required for consolidation of a new memory, a reminder of the experience that generates that memory involves only local synthesis at the synapse.

Unequivocal support for this hypothesis would require a drug that specifically blocks synaptic/dendritic protein synthesis, but spares somatic protein synthesis, and preferably, the identi-
fication of the proteins involved; in the absence of such proof, our finding may be open to alternative interpretations—for instance, that the Ani-induced amnesia results not directly from Ani’s effect as a protein synthesis inhibitor, but from some other or downstream consequence—such as a change in local levels of free amino acids as a result of inhibiting synthesis. However, what is now clear is that in both this avian aversive learning paradigm and in some mammalian ones, as well as in the experience of a delay between the initial experience and the reminder (Anokhin et al. 2004). The circuitry involved also appears to differ, as a reminder enhances metabolic activity in the MS but not the IMMP, the region involved in the post-training cascade (Salinska et al. 2002). The issue of whether and under what conditions the amnesia is permanent rather than transient is induced by antisense oligos to it (Lee et al. 2004). Further, the transcription factor Zif268 is not required for initial consolidation, but is apparently engaged following a reminder as amnesia transients from the events following training. In mammalian systems, there are differences in the types of immediate early genes activated (Taubenfeld et al. 2001; Izquierdo and Cammarota 2004), while amygdalar circuits involved in consolidation are apparently not required for reconsolidation (Bahar et al. 2004). In the chick, the dose dependency of amnesia to agents such as Ani or 2-deoxyglucose, and the temporal dynamics of the amnestic response, differ between training and reminder (Anokhin et al. 2002). The machinery involved also appears to differ, as a reminder enhances metabolic activity in the MS but not the IMMP, the region involved in the post-training cascade (Salinska et al. 2004).

The pharmacological lability of memory following a reminder is far from a universal phenomenon, and often seems to involve different molecular processes from those following initial training. Thus, in contextual fear conditioning in the rat, the transcription factor Zif268 is not required for initial consolidation, but is apparently engaged following a reminder as amnesia is induced by antisenol oligos to it (Lee et al. 2004). Furthermore, the effects of inhibitors appear to vary with species and task. For instance, in contrast to retrieval after a reminder for aversive learning tasks such as fear conditioning, passive or inhibitory avoidance or conditioned taste-aversion instrumental responses are not affected by protein synthesis inhibitors following a reminder (Hernandez and Kelley 2004). Yet, in a rewarding odor-discrimination task in rats (Torras-Garcia et al. 2005), as with passive avoidance in chicks (Litvin and Anokhin 2000), NMDA receptor blockade following reminder is amnestic. The effects of such inhibitors following aversive tasks is dependent on the interval between the initial experience and the reminder (Anokhin et al. 2002; Alberini 2005), and there is still a conflict of evidence over whether and under what conditions the amnesia is permanent rather than transient. The results presented in this study also require us to reconsider the nature of the biochemical processes initiated by the reminder, suggesting as they do that, at least in this form of avian learning, engage local synaptic and dendritic processes, not necessarily in the same cells or brain regions as those involved in initial consolidation. This distinction may account for the transient nature of the amnesia produced by protein synthesis inhibitors following a reminder in this experimental paradigm.

## Materials and Methods

### Animals and training

Ross Chunky chicks were hatched and reared in our own incubators. Day-old chicks were placed in pairs in pens, pre-trained, and trained as described previously (Loxner and Rose 1983). Briefly following a stabilization period of 1 h in the pens, the chicks were pre-trained by three presentations of a small (2.5 mm diameter) white bead over a 10-min period. Chicks that pecked at least twice during pre-training (>85%) were then presented with a 4-mm diameter chrome bead dipped in the bitter-tasting methylantranilate for 20 sec. Chicks that pecked this bead and evinced a disgust response (>90%) were considered to be trained. At appropriate times, later chicks were tested by being offered the dry chrome bead for 30 sec and 5 min later, the pre-training white bead. Chicks that avoided the white on test were considered to have retained the memory for the aversive experience and to be able to discriminate. Retention was calculated as the percent in each group that showed avoidance and discrimination. Each chick was trained and tested only once. The differences between hatches were tested with ANOVA (Single factor analysis). Statistical analysis of behavioral data was by analysis of variance (ANOVA) and where significant differences were found between groups, further analysis was made using a post hoc least-significant difference test.

### Drugs

Chicks were injected at different time points pre- or post-training into the left and right IMMP or MS. Bilateral intracranial injections were made using a custom-built head holder (Davis et al. 1982) and a 5-µL Hamilton syringe fitted with a plastic sleeve to allow appropriate penetration. Correct placement was ensured by and was routinely visually monitored post-mortem. Anisomycin (ANI; Sigma) was dissolved in sterile 3 M HCl made in saline solution and the pH adjusted to 7.4 with 3 M NaOH. Colchicine (Sigma) was dissolved directly in sterile saline. The total dose of Ani was 125 nmol/hemisphere, and for Colchicine, 15 nmol/hemisphere. The volume of injected drugs was 2 µL.

### Protein synthesis and axonal flow

Chicks were injected as above with 14C-leucine (L-[14C(U)], 0.0018 MBq/hemisphere) (ARC65), followed 15 min later by injection of Colchicine (15 nmol/hemisphere) (group LC). Controls received the same amount of 14C-leucine (group L). n = 5 for each group. Three hours later, birds were decapitated and IMMP samples were dissected out and homogenized in an ice-cold buffered sucrose solution (1:10 w/v, 5mM HEPES at pH 7.4, 0.52 M sucrose) supplemented with protease inhibitor cocktail (Roche, Diagnostic) for synaptoneurosomal preparation (Oestreicher and van Leeuwen 1975). The total homogenate was centrifuged for 10 min at 1000g. The supernatant was enriched in synaptoneurosomal preparation (Oestreicher and van Leeuwen 1975). The total homogenate was centrifuged for 10 min at 1000g. The supernatant was centrifuged for 45 min at 17,000g to give a crude synaptoneurosomal and mitochondriaic pellet. The obtained pellet was resuspended in ice-cold buffered sucrose and spun over a 1.2 M sucrose cushion to remove remains of myelin and larger mitochondria. The final pellet was enriched in synaptoneurosomal (see Fig. 2D). Aliquots were taken at each step throughout the procedure for protein Bradford (1976) and for total and incorporated 14C-leucine measurement (see Fig. 2E). Specific radioactivity (dpm/mg protein) was determined by trichloroacetic acid precipitation.

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