Memory decay and susceptibility to amnesia dissociate punishment- from relief-learning

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Painful events shape future behaviour in two ways: stimuli associated with pain onset subsequently support learned avoidance (i.e. punishment-learning) because they signal future, upcoming pain. Stimuli associated with pain offset in turn signal relief and later on support learned approach (i.e. relief-learning). The relative strengths of such punishment- and relief-learning can be crucial for the adaptive organization of behaviour in the aftermath of painful events. Using Drosophila, we compare punishment- and relief-memories in terms of their temporal decay and sensitivity to retrograde amnesia. During the first 75 min following training, relief-memory is stable, whereas punishment-memory decays to half of the initial score. By 24 h after training, however, relief-memory is lost, whereas a third of punishment-memory scores still remain. In accordance with such rapid temporal decay from 75 min on, retrograde amnesia erases relief-memory but leaves a half of punishment-memory scores intact. These findings suggest differential mechanistic bases for punishment- and relief-memory, thus offering possibilities for separately interfering with either of them.

1. Introduction

A painful event has two sides: A ‘negative’ aspect at its onset and a ‘positive’ aspect at its offset, at the moment of relief [1]. Flies, for example, avoid an odour once it has been associated with the onset of an electric shock (odour → shock; henceforth called punishment-learning); yet they approach an odour once it has been associated with shock offset (shock → odour; henceforth called relief-learning; [2,3]). Similar Janus-headed results are found in rats and man: visual cues associated with shock onset potentiate startle, whereas cues associated with shock offset attenuate startle [4,5]. Thus, if embedded into a natural string of events, the net effect of such an adverse life event may depend on the relative strengths of oppositely valenced memories related to its onset (punishment-memory) and its offset (relief-memory). Here, we compare the time course of decay, as well as the susceptibility to post-training cold-amnesia, between these two kinds of memory. The differences we find in both these parameters suggest a dissociation of the mechanisms underlying punishment- and relief-memories.

2. Material and methods

Training and testing followed standard methods with the modifications described in detail in Yarali et al. [3] and electronic supplementary material, figure S1. Please note...
training trials were concatenated in both cases (for details, see the whereas for relief-training it was presented upon shock offset; six memory is obviously much stronger than relief-memory, as ferring flies into ice-cold plastic vials and then keeping them on 120 min after training. Cold-amnesia was implemented by trans- and then either did or did not receive cold-amnesia at 60 min such that flies underwent either punishment- or relief-training, retention period).

These vials either remained within the experimental room yet outside the conditioning apparatus (50, 75 and 240 min retention periods) or were transferred overnight to the culture facility (24 h retention period).

(a) Memory decay
This experiment used a 2 × 5 experimental design (figure 1a), such that flies underwent either punishment- or relief-training, and were then tested after retention periods of 25, 50, 75, 240 min, or 24 h. During the retention period and in the intervals between training trials, flies were kept in their regular food vials. These vials either remained within the experimental room yet outside the conditioning apparatus (50, 75 and 240 min retention periods) or were transferred overnight to the culture facility (24 h retention period).

(b) Resistance to cold-amnesia
This experiment used a 2 × 2 experimental design (figure 2a), such that flies underwent either punishment- or relief-training, and then either did or did not receive cold-amnesia at 60 min after training. Testing then took place 60 min later, i.e. at 120 min after training. Cold-amnesia was implemented by transferring flies into ice-cold plastic vials and then keeping them on ice for 2 min before transferring them back to regular food vials.

(c) Statistics
Non-parametric statistics were used throughout. Kruskal–Wallis (KW) tests and Mann–Whitney U (MWU) tests were used to compare LI values between multiple and two groups of flies, respectively. One-sample sign tests (OSS) were used to determine whether scores of a given group were significantly different from zero. Throughout, a significance level of \( p < 0.05 \) was adopted. For multiple comparisons within a dataset, critical \( p \)-levels were adjusted by a Bonferroni correction (\( p < 0.05 \) divided by the number of comparisons) to maintain the experiment-wide error rate at 5 per cent. Data are plotted as box plots, representing the median as the middle line, the 25 and 75 per cent quantiles as boundaries of the box and the 10 and 90 per cent quantiles as whiskers.

3. Results
(a) Memory decay
In terms of absolute learning scores (figure 1b), punishment-memory is obviously much stronger than relief-memory, as has been reported previously [2,3]. For the current context, it is important that punishment-memory scores decay as the retention period lengthens (KW: \( p < 0.05 \), \( N = 20 \) in all cases, d.f. = 4, \( H = 36.69 \); MWU comparing later retention periods to the earliest one (25 min): \( p < 0.05 \) in all cases, \( U = 97.00, 57.00, 62.00, 12.00 \)). In fact, within 75 min punishment-memory has decayed to approximately 50 per cent of the initial score. Critically, however, punishment-memory remains detectable for at least 24 h (OSS for each retention period: \( p < 0.05/5 \) in all cases).

Relief-memory scores also change with lengthening of the retention period (KW: \( p < 0.05 \); \( N = 51, 35, 46, 43, 40 \); d.f. = 4, \( H = 13.82 \), but in a different way from punishment-memory scores. That is, compared with the shortest retention interval of 25 min, relief-memory remains stable during at least the first 240 min (MWU: \( p > 0.05/4 \) in all cases, \( U = 839.00, 1161.00, 833.00 \); mind the tendential difference for the 240 min retention period). At 24 h after training, however, relief-memory scores have decayed significantly (MWU: \( p < 0.05/4 \); \( U = 634.5 \)). Fittingly, relief-learning memory scores are significantly positive only for retention periods of up to 75 min (OSS: \( p < 0.05/5 \) in all cases except for 240 min, where \( p = 0.014 \) and 24 h, where \( p = 0.4 \)).
4. Discussion

We provide the first systematic comparison of the temporal dynamics of punishment- versus relief-memory. Over the first 4 h following training, relief-memory decays much slower than punishment-memory (figure 1b). This slow decay is reminiscent of the slow initial decay rate of sugar reward-memory [6–8]. With respect to longer retention periods, however, relief-memory differs from both punishment- and reward-memories: multiple, temporally spaced training trials result in detectable punishment-memory beyond 24 h [9,10], whereas for such long-term reward-memory, a single training trial suffices [8,11]. For relief-learning, despite using multiple, spaced training trials, we find no appreciable memory scores at 24 h (figure 1b). Thus, the temporal pattern of decay for relief-memories differs from both punishment- and reward-memories.

Regarding cold-amnesia, both punishment- and reward-memories are only partially susceptible within the first 2 h following training [6–10,12–14]. That is, cold-amnesia typically spares a so-called amnesia-resistant component of reward- and of punishment-memory. Indeed, we confirm that punishment-memory 1 h after training is composed of an amnesia-sensitive component and an amnesia-resistant component (figure 2). Critically, however, cold-amnesia abolishes relief-memory completely (figure 2). Given that for punishment-memory, anaesthesia-sensitive versus -resistant components of memory seem to have partially different genetic requirements (for proper function of e.g. the amnesiac and rutabaga [10], synapsin [13] and bruchpilot [14] genes), it would be interesting to look for roles of these genes in relief-learning.

If, within a single subject, the events before and after a traumatic episode were to induce punishment- and relief-memory, our finding that both these forms of memory differ in strength and susceptibility to retrograde amnesia may be of practical importance: while trying to erase punishment-memory, one may unwittingly also erase relief-memory. Dependent on the relative strength of these memories and the relative effectiveness of the treatment, the net effect of such manipulation may make the overall-mnemonic effect of the traumatic episode even more adverse.

References


