Evidence suggests that extinction is new learning. Memory acquisition involves both short-term memory (STM) and long-term memory (LTM) components; however, few studies have examined early phases of extinction retention. Retention of auditory fear extinction was examined at various time points. Shortly (1–4 h) after extinction acquisition spontaneous recovery was high compared to that after longer delays (8–24 h). Recall of a consolidated extinction trace was also impaired if it was preceded 1 h by extinction of a novel CS; propranolol did not attenuate this effect. These results suggest poor extinction retention reflects a retrieval impairment caused by the aversive experience of extinction training.

In Pavlovian fear conditioning pairing a neutral stimulus with an unconditioned aversive stimulus (US) results in conditioned learning, whereby the previously neutral stimulus comes to elicit fear responses. Conversely, the conditioned stimulus (CS) repeatedly presented in the absence of the US leads to reduction of the fear behavior known as extinction (Pavlov 1927). Although there has been some debate over the exact nature of extinction, it is widely accepted that this phenomenon is new learning rather than unlearning of the original association (Rescorla 1997; Bouton 2004). Return of the extinguished behavior is observed in tests of spontaneous recovery (Pavlov 1927), renewal (Bouton and King 1983), and reinstatement (Rescorla and Heth 1975; Bouton and Bolles 1979).

Both behavioral (McGaugh 1966) and molecular (Kandel 2001) studies suggest that following learning there are two distinct phases of memory: short-term memory (STM) and long-term memory (LTM). STM, lasting on the order of minutes to hours, involves covalent modifications of preexisting proteins, while LTM is much less transient and involves protein synthesis, new gene expression, and changes in synaptic structure. Disruption of protein synthesis following learning leads to impairments in LTM retention while sparing STM (Davis and Squire 1984). STM for fear extinction is often considered to be the within-session responding during extinction training, whereas LTM for fear extinction is measured by successful extinction recall usually tested upward of 24 h (Quirk 2002). Tests of long-term extinction memory are frequently used to measure spontaneous recovery, the return of the original trace that occurs with the passage of time; more recovery is observed with increasing delays between extinction and test (Pavlov 1927; Elson 1939; Burdick and James 1970; Robbins 1990). Surprisingly, few studies have tested memory for fear extinction within 24 h of extinction acquisition.

Berman et al. (2003) explored the course of an extinguished conditioned taste aversion (CTA) memory, testing retention of extinction at intervals as short as 30 min following extinction. They found that the extinguished behavior was not evident shortly after extinction but became apparent with a delay of 2 h. The authors further determined that the absence of extinction shortly after extinction training was likely due to generation of a short-term aversive trace that blocked the immediate development of extinction. These results are inconsistent with the idea that recovery of the original memory increases with time. Further, it remains unclear whether extinction of an auditory fear memory will exhibit a similar pattern of recovery. Here we addressed this question and further attempted to determine a mechanism mediating our findings.

Adult male Sprague-Dawley rats (Charles River Laboratories, Quebec) initially weighing 300–325 g were pair housed in plastic Nalgene cages. They were maintained on a 12-h light/dark cycle (lights on at 7 a.m., off at 7 p.m.) with food and water available ad libitum. All procedures were in accordance with the Canadian Council on Animal Care guidelines and were approved by the McGill University Animal Care and Use Committee. Two conditioning contexts were used: Context A for fear conditioning and Context B for extinction and testing. Varying dimensions, such as shape, illumination, scent, visual and tactile features, and ambient sound, minimized generalization between contexts.

We first determined whether extinction of an auditory fear memory is expressed shortly after extinction acquisition by comparing levels of freezing to a tone at different time points following extinction. Rats were divided into four groups and conditioned using two tone presentations co-terminating with shock (0.8 mA, 0.5-sec shock; 5-kHz, 75-dB, 20-sec tone). Three days later all groups received 16 non-reinforced stimulus presentations over 30 min in Context B. Memory for extinction was tested in Context B 1 h, 4 h, 8 h, or 24 h after extinction training. Extinction produced roughly equivalent within-session reductions in freezing. Figure 1B shows that recovery was greater in groups tested 1 h and 4 h after extinction than the groups tested at 8 h and 24 h. A 4 (Group) × 2 (Block) mixed ANOVA comparing the last two-trial block of extinction with the test confirmed a main effect of Group, $F_{(3, 45)}= 5.41, \ P < 0.05$, and a main effect of Block, $F_{(1, 45)}= 50.54, \ P < 0.001$. There was also a Group × Block interaction, $F_{(3, 45)}= 4.65, \ P < 0.05$, indicating that freezing was higher in groups tested after a short delay. Overall, the results suggest that extinction behavior is absent at short-term time points after extinction acquisition but develops with longer delays between extinction training and test. Consistent with this, in rats that had received two spaced extinction sessions, an extinction-test interval of 2 min resulted in more recovery of freezing to a feared context than an interval of 24 h (Morris et al. 2005a).

The transient recovery may reflect a breakdown between phases of memory that support performance (i.e., intermediate-term memory) or a retrieval impairment due to physiological fluctuations produced by the extinction session. In a second experiment we tested whether the absence of extinction behavior was...
a specific property of the CS being extinguished or would generalize to other previously extinguished CSs. Animals were fear conditioned to both a tone and a white noise (2-kHz, 75-dB, 20-sec hiss) during separate sessions on consecutive days, the presentation order was counterbalanced such that half the animals received tone as CS1 and white noise as CS2. Both stimuli were then extinguished in separate extinction sessions on successive days. Expression of extinction of CS1 was tested either 1 h or 24 h after extinction acquisition, respectively. Rats in the short-delay groups (1 h and 4 h) exhibited high levels of freezing compared to animals tested 1 h after last block of extinction and to the longer delay testing groups (\(t^*\) \(P < 0.05\)).

These results were similar to those of the first experiment: a test given shortly after extinction led to fear recovery. However, in this experiment return of responding to the CS was not a result of a deficit in short-term or intermediate-term extinction memory. Indeed, expression of extinction may be impeded by changes in the physiological state of the animal that occur during extinction training.

Adrenergic transmission has been implicated in the formation and persistence of emotional memories (Izquierdo and Medina 1995; Cahill et al. 2000). Similarly, propranolol, a \(\beta\)-adrenergic receptor antagonist, has been used to reduce fear and anxiety in both humans and rats (Gorman and Dunn 1993; Kent et al. 2002). Morris et al. (2005b) discovered that a brief exposure to a feared context was sufficient to reinstate freezing to an extinguished CS, but only if the exposure occurred 2 min before the CS test and not 24 h prior. The reinstatement of freezing following exposure to a feared context appears to be dependent on \(\beta\)-adrenergic activation as treatment with propranolol attenuated this effect.

In a third experiment we set out to address whether adrenergic activity was mediating the transient recovery from extinction. Behavioral procedures of Experiment 3 were identical to those of Experiment 1; however, 20 min prior to the CS1 retention test, animals were injected intraperitoneally with 5 mg/kg of propranolol or vehicle (Fig. 3A). All results are shown in Figure 3B. A three-way mixed ANOVA of Test (1 h vs. 24 h) \(\times\) Drug (Prop vs. Veh) \(\times\) Block (last extinction vs. test) revealed no main effect of drug or testing time and no interaction between the two. Analysis did indicate a significant main effect of Block, \(F_{(3,28)} = 19.38, P < 0.0001\), and significant interaction of Block \(\times\) Test, \(F_{(3,28)} = 14.10, P < 0.001\). Overall, these results indicate that under these conditions treatment with propranolol was unable to prevent the transient recovery observed shortly after extinction.

This is in contrast to previous findings in which propranolol was able to block reinstatement of extinguished fear following exposure to a dangerous context (Morris et al. 2005b). Nonetheless, in these experiments propranolol was given prior to the context exposure which reinstated freezing to an extinguished CS. Consequently, we determined whether propranolol given before extinction was able to block increases in adrenergic activity that might develop during extinction, which would affect the short interval retention test. Examining only the 1-h group, propranolol was injected 20 min prior to extinction acquisition (Fig. 3C). Results indicated no differences between treatment groups as...
both groups showed significant recovery during the 1-h retention test after extinction acquisition as indicated by a significant effect of block, \( F_{1,14} = 4.59, P < 0.0001 \). Even at this time point propranolol was unable to prevent recovery of the original fear conditioning.

The present experiments suggest that memory for auditory fear extinction does not have the same temporal pattern of behavioral expression as predicted by our current understanding of spontaneous recovery. Although significant within-session (short-term) extinction was observed, memory for extinction was absent at 1-h and 4-h time points. Good extinction retention was again observed in the 8-h and 24-h test groups. This is consistent with the profile of behavior following CTA extinction (Berman et al. 2003) and context fear extinction (Morris et al. 2005a). Generally, the amount of spontaneous recovery is considered a function of the time between extinction and test, such that greater recovery is observed with longer retention intervals. The present results suggest that spontaneous recovery follows a nonmonotonic function that resembles the U-shaped retention curve first described by Kamin (1957).

The “Kamin effect” has been widely demonstrated (Gerber and Menzel 2000; Sutton et al. 2001; Rudy and Wright-Hardest 2005; McNally et al. 2008). It was initially thought to be a result of two memory systems. One system was responsible for immediate retention, losing strength rapidly after acquisition. A second system, requiring time for consolidation, gradually assumed responsibility for memory retention and expression. Any transient lapse in memory retention was thought to reflect a discontinuity between these two systems. A second interpretation has also been proposed suggesting that the observed lapse in memory is a result of a retrieval impairment due to a discrepancy in the internal state of the animal at intermediate intervals compared to immediate or long-term intervals (Klein and Spear 1970; Klein 1972).

Discontinuous expression of extinction may reflect absence of the memory or the inability to retrieve the extinction memory and respond accordingly. Berman et al. (2003) note that extinction itself may be aversive and lead to hormonal and neurochemical changes that promote reinstatement of fear. Indeed, we found that testing a previously extinguished CS (CS1) 1 h after extinction of a second CS (CS2) elicited spontaneous recovery. If, however, the CS1 extinction retention test occurred 24 h after extinction of CS2, there was little recovery. The absence of extinction was not specific to a recently extinguished CS and instead generalized to a previously extinguished CS; thus, the aversive experience of extinction may be sufficient to reinduce freezing to a different previously extinguished CS. These results are consistent with the observations that a recent pre-exposure to a feared context can reinstate extinguished auditory fear memory (Morris et al. 2005a) and exposure to a different malaise associated taste is sufficient to reinduce an extinguished conditioned taste (Berman et al. 2003).

Interestingly, in an appetitive task in which the CS is not an aversive stimulus, memory for extinction is observed 5 h following extinction training (Brooks and Bouton 1993). Thus, the “lapse” of extinction at a short retention interval may not be a characteristic of all extinction memories. Also contrary to our findings, Quirk (2002) showed that 30 min after extinction of an auditory fear memory, extinction retention was successful. Discrepancies in the results may be due to differences in behavioral protocols or they may indicate that impaired extinction recall does not occur until after 30 min.

It is well established that context can influence responding to a CS after extinction (Bouton et al. 2006). Beyond the modulatory effect of the physical context, there is evidence that the emotional state of the animal can serve as a “context” and lead to the return of the fear responding (Richardson et al. 1984). It has previously been shown that administration of propranolol prior to a brief period spent in a feared context prevents reinstatement of freezing to an extinguished CS that would have otherwise occurred (Morris et al. 2005b). Based on this finding, we hypothesized that the adrenergic system, in response to the aversive experience of being reexposed to the feared CS during extinction, might be mediating the transient spontaneous recovery. Propranolol was administered 20 min prior to the extinction session and then tested for retention at the 1-h time point. Under both conditions propranolol was unable to prevent the transient recovery of freezing observed in the 1-h group. These results suggest that blocking β-adrenergic activity is unable to attenuate the effects of emotional arousal resulting from the
recent experience of extinction. This is in contrast to previous findings in which propranolol was able to block reinstatement of extinguished fear following exposure to a dangerous context (Morris et al. 2005b).

The currently established neurocircuitry for fear extinction memories involves the basolateral amygdala (BLA), the central amygdala (CeA), the ventral medial prefrontal cortex (vmPFC), and the hippocampus (HC). The infralimbic cortex (IL) appears to be the critical site in the vmPFC involved in extinction consolidation and retrieval (Laurent and Westbrook 2009; Sierra-Mercado et al. 2011). Inputs to the intercalated (ITC) cells in the amygdala from the IL are also necessary for extinction (Likhith et al. 2008) and activity in the ITC subsequently suppresses activity in CeA neurons (Quirk et al. 2003; Amano et al. 2010). Given that prelimbic (PL) activity impairs extinction and increases fear expression (Vidal-Gonzalez et al. 2006) while IL activity suppresses fear after extinction (Quirk et al. 2006), it is reasonable to suspect the transient recovery from extinction we observe soon after acquisition may result from disruptions in IL activity and ITC projections to the CeA necessary for fear suppression.

Exposure to stressors results in a number of neurochemical changes within the mPFC. Extinction of eyelid conditioning causes an increase in the extracellular levels of dopamine and norepinephrine in the mPFC that decrease slowly in the 2 h following (Hugues et al. 2007). Either of these in addition to other neuromodulators may be involved in inhibiting activity in the IL. Further determining the neurochemical changes that accompany IL activation will help elucidate the mechanisms involved in the transient recovery from extinction.

In conclusion, our results suggest that auditory fear extinction is not expressed shortly after extinction training. Poor extinction recall, as measured by spontaneous recovery, follows the nonmonotonic function characteristic of the “Kamin effect.” This impairment in retention does not appear to be a result of a storage failure, nor is it mediated solely by β-adrenergic activity. Understanding the mechanisms and neural circuitry involved in the formation and expression of extinction remains to be fully understood. Future studies directed in this area will likely shed light on the interplay between the expression of fear and the expression of extinction and may account for the transient recovery from extinction.

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Temporal dynamics of recovery from extinction shortly after extinction acquisition

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