Shifting from fear to safety through deconditioning-update

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Abstract Aversive memories are at the heart of psychiatric disorders such as phobias and post-traumatic stress disorder (PTSD). Here, we present a new behavioral approach in rats that robustly attenuates aversive memories. This method consists of ‘deconditioning’ animals previously trained to associate a tone with a strong footshock by replacing it with a much weaker one during memory retrieval. Our results indicate that deconditioning-update is more effective than traditional extinction in reducing fear responses; moreover, such effects are long lasting and resistant to renewal and spontaneous recovery. Remarkably, this strategy overcame important boundary conditions for memory updating, such as remote or very strong traumatic memories. The same beneficial effect was found in other types of fear-related memories. Deconditioning was mediated by L-type voltage-gated calcium channels and is consistent with computational accounts of mismatch-induced memory updating. Our results suggest that shifting from fear to safety through deconditioning-update is a promising approach to attenuate traumatic memories.

Introduction

Memory is a dynamic process that allows for adaptation to the demands of a continuously changing environment. The ability to update old memories in accordance with new experiences is crucial for maintaining their relevance over time. Particularly, it has been shown that after retrieval (or reactivation), memories may undergo a cycle of labilization and restabilization commonly known as reconsolidation (Nader et al., 2000). The labile state induced by this process can thus allow changes in memory strength and/or content (De Oliveira Alvares et al., 2013). This has been most extensively studied in aversive conditioning paradigms in rodents and humans and is of potential relevance to the management of psychiatric disorders involving dysfunctional memories (Monfils and Holmes, 2018).

Reconsolidation has been described in several experimental paradigms and species, from invertebrates to humans, suggesting that it might be a fundamental property of memories (Nader and Einarsson, 2010). Beyond its biological role in memory updating, it opens a window of opportunity...
as a potential therapeutic strategy to modify pathological memories. Several studies in the last decades have attempted to pharmacologically or behaviorally disrupt the reconsolidation of traumatic memories (Beckers and Kindt, 2017), as at least in some situations, this strategy can be less susceptible to spontaneous or induced recovery than traditional extinction (Duvarci and Nader, 2004; Monfils et al., 2009). However, most treatments that interfere with memory reconsolidation are toxic and cannot be readily administered to humans. Thus, in spite of almost two decades of research on memory reconsolidation, the evidence for the efficacy of reconsolidation-blocking treatments in clinical settings is still limited (Monfils and Holmes, 2018).

Research on reconsolidation-extinction boundaries suggests that the transition from one process to the other depends on the degree of mismatch between the original memory and the reactivation experience. Many studies have suggested that some degree of mismatch or prediction error is necessary for reconsolidation to occur (Lee, 2009; Fernández et al., 2016); however, if prediction error is too high, extinction may occur instead (Suzuki et al., 2004; Sevenster et al., 2014). Computational models suggest that, in models in which prediction error is low, memory updating/reconsolidation mechanisms are preferentially engaged, as the new experience is recognized as a new instance of the former one; however, as mismatch increases, the chances of it being attributed to a new latent cause (Gershman et al., 2017) or forming a new attractor in a neural network (Osan et al., 2011) increases.

This framework suggests that lowering the degree of mismatch between learning and reexposure might conceivably potentiate memory updating during extinction. This has been the rationale behind so-called retrieval-extinction procedures (Monfils et al., 2009; Kredlow et al., 2016) and has also been explored in short-term extinction protocols (Gershman et al., 2013). In this work, we propose a novel approach for long-term attenuation of traumatic memories that we term ‘deconditioning-update’. This strategy consists in weakening fear memories by updating the aversive information, substituting the original US by a mildly aversive stimulus during the plastic state induced by reactivation.

Results
In order to approach our hypothesis, male Wistar rats were trained in auditory fear conditioning, where they received five conditioning trial tones (CS) that co-terminated with a 0.5-mA, 1-s footshock (US). On days 3 to 6 (reactivation), animals received 3 CSs during a 400-s daily session in a different context. In the no-footshock group, CSs were presented in the absence of shock, while in the deconditioning-update group, each tone co-terminated with a 0.1-mA footshock. A third group remained in the homecage (control group). On day 7, both groups were tested in the reactivation context with 3 CSs. On day 8, animals were tested in the training context for renewal, and, on day 28, they were retested for spontaneous recovery (Figure 1A).

Over the course of reactivation sessions, both groups presented a decrease in freezing, but this was more marked in the deconditioning-update group (Figure 1B), which presented a decrease in freezing of around 70% in comparison to the no-footshock group and 80% in comparison to the homecage control group in the test session (Figure 1C). Animals in the deconditioning-update group also had lower freezing responses in the renewal (Figure 1D) and spontaneous recovery (Figure 1E) tests, although it should be noted that memory recovery was not observed in the no-footshock group either, perhaps due to a ceiling effect caused by insufficient extinction (for complete statistics, see Supplementary file 1). Pre-CS freezing varied between 37% and 69% in both groups throughout the extinction sessions (Supplementary file 12), suggesting some degree of generalization between both contexts.

When performing the same experiment, but with each tone co-terminating with a 0.3-mA footshock in the reactivations, no fear reduction occurred (Figure 1—figure supplement 1); on the contrary, the footshock group presented higher levels of freezing than the no-footshock group throughout the reactivation sessions, as well as in the test. This result is in accordance with a recent study showing that fear memory may be strengthened by reactivation with a 0.3-mA footshock (Ferrara et al., 2019a). It also rules out that the suppression of freezing during reactivations by deconditioning-update may be due to inhibition of delay (i.e. animals learning to freeze only at the end of the CS with extended practice). When using a single reactivation session with 0.1mA, no
Figure 1. Weakening fear memory through deconditioning-update training. (A) Experimental design: male rats were fear-conditioned with five tone-shock pairings (context A; 5 CS + US, 0.5mA). 48 hr later, the no-footshock and footshock (deconditioning-update) groups were exposed to four daily reactivation sessions (context B). After this, animals underwent test (context B), renewal (context A) and spontaneous recovery (context B) sessions. Black circles represent context A, while white rectangles represent context B. (B) Freezing levels during reactivation sessions. Rats exposed to weak footshocks during reactivation sessions showed a significant reduction in freezing responses, maintained during the test (C), renewal (E) and spontaneous recovery (D) sessions. (F) Experimental design: female rats were fear-conditioned (context A; 5CS+US, 0.5mA). 48 hr later, the no-footshock and footshock groups were exposed to four daily reactivation sessions (context B). After this, all groups underwent test, renewal, renewal (context A) and spontaneous recovery (context B) sessions. (G) Reactivations

% Freezing

Day 3 Day 4 Day 5 Day 6

No Footshock Footshock

(H) Test

% Freezing

Control Footshock No Footshock

(I) Renewal

% Freezing

Control Footshock No Footshock

(J) Spontaneous Recovery

% Freezing

Control Footshock No Footshock

(K) Retraining

% Freezing

Control Footshock No Footshock

Figure 1 continued on next page
and spontaneous recovery sessions. Animals were reconditioned (context A; 3CS+US, 0.5mA) on the next day and retested 24 hr later. (G) Freezing levels during memory reactivation. Rats exposed to weak footshocks showed a significant reduction in freezing responses, maintained during the test (H), renewal (I), spontaneous recovery (J) and retraining test (K) sessions. Bars represent mean ± SEM. Statistical comparisons were performed using two-way repeated-measures ANOVA followed by a Bonferroni post-hoc (reactivation sessions) or one-way ANOVA followed by Tukey post-hoc (test, renewal, spontaneous recovery, and retraining test). *p<0.05, **p<0.005, ***p<0.0005, ****p<0.0001. For full statistics, see Supplementary file 1. For pre-CS freezing values, see Supplementary file 12.

The online version of this article includes the following source data and figure supplement(s) for figure 1:

Source data 1. Raw data of Figure 1.

Figure supplement 1. Deconditioning-update does not occur with 0.3-mA shocks.

Figure supplement 2. A single reactivation session does not update fear memory.

Figure supplement 3. Deconditioning-update is not due to US devaluation.

Figure supplement 4. Deconditioning-update does not occur with unpaired shocks in the reactivation sessions.

difference was found in the test in comparison to the no-footshock group (Figure 1—figure supplement 2), suggesting that deconditioning-update requires several sessions to take place.

One could argue that the exposure to weak footshocks could simply lead to habituation and consequent devaluation of the US, without altering the conditioned association itself (Rescorla, 1973; Rescorla and Heth, 1975). In order to test this alternative interpretation, rats were conditioned as described above and the same 3 0.1-mA weak footshocks were given in another context without being paired with the tones (Figure 1—figure supplement 3A). In this case, no fear reduction was found in comparison to homecage controls. (Figure 1—figure supplement 3B). To further rule out the devaluation hypothesis, another set of animals was submitted to reinstatement after deconditioning, in order to test whether restoring the original footshock valence outside of the extinction context might lead to memory recovery. We found that the deconditioning group expressed a lower fear level when compared with the no-footshock group in the test even after reinstatement (Figure 1—figure supplement 3E), suggesting that the deconditioning-update effect is not due to US devaluation, but rather to updating of the CS-US association.

In an additional experiment, we investigated whether the stress induced by weak footshocks during reactivation could be enhancing extinction by itself. Animals underwent either the deconditioning-update protocol described in Figure 1A, or a similar protocol in which both tones and footshocks were applied during the reactivation sessions, but with no pairing between them. We found that this non-contingent tone-footshock protocol did not attenuate fear expression, leading to higher freezing than deconditioning-update during reactivations and in the test (Figure 1—figure supplement 4).

Next, we asked whether the deconditioning-update effect would also be efficient in reducing fear in female rats. As observed in males, females from the deconditioning-update group showed lower freezing levels throughout the reactivation sessions, as well as in the test session, when compared with the control group (Figure 1H). The same pattern was maintained in the renewal and spontaneous recovery tests (Figure 1I and J, respectively). As a further way to test for persistence of the original memory, we performed a retraining session in the original context with 3 0.5-mA CS-US pairings 24 hr after the spontaneous recovery test to assess savings. The deconditioning-update group had lower freezing compared with the other groups in a subsequent test session in the extinction context, suggesting that our protocol also lowers the rate of reacquisition of an aversive memory (Figure 1K).

Boundary conditions such as training intensity and memory age have been reported to prevent fear memories from being modified. Protocols with high training intensity make memory less prone to interference by pharmacological agents in the reconsolidation window (Frankland et al., 2006; Suzuki et al., 2004; Wang et al., 2009), while older memories are also less susceptible to modification than recent ones (Milekic and Alberini, 2002; Frankland et al., 2006; Suzuki et al., 2004; Bustos et al., 2009). Thus, our next experiments investigated whether deconditioning-update could attenuate fear expression in these cases. First, we trained the animals in the same protocol described above, but starting reactivations 40 days after conditioning (Figure 2A). Once more, the
Figure 2. Deconditioning-update weakens both remote and strong fear memories. (A) Experimental design for remote memory: rats were fear-conditioned with five tone-shock pairings (context A; 5 CS + US, 0.5mA). Starting 40 days later, the no-footshock and footshock (deconditioning-update) groups were exposed to daily reactivation sessions (context B). Then, all groups underwent test (context B), renewal (context A) and spontaneous recovery sessions (context B).
Figure 2 continued

(context B) sessions. Black circles represent context A, while white rectangles represent context B. (B) Freezing levels during reactivation sessions. Rats exposed to weak footshocks during reactivation sessions showed similar freezing levels to no-footshock animals during the test session (C) and lower freezing levels at the renewal (D) and spontaneous recovery (E) ones. (F) Experimental design for strong training (5CS+US, 1mA). (G) Freezing levels during reactivation sessions. Rats exposed to weak footshocks during reactivation sessions showed a significant reduction in freezing responses that was maintained during the test (H), renewal (I) and spontaneous recovery (J) sessions. Bars represent mean ± SEM. Statistical comparisons are performed using two-way repeated-measures ANOVA followed by a Bonferroni post-hoc (reactivation sessions) or one-way ANOVA followed by a Tukey post-hoc (test and spontaneous recovery). *p<0.05; **p<0.005; ***p<0.0005; ****p<0.0001. For full statistics, see Supplementary file 2. For pre-CS freezing, see Supplementary file 13.

The online version of this article includes the following source data and figure supplement(s) for figure 2:

Source data 1. Raw data of Figure 2.
Figure supplement 1. Deconditioning-update weakens strong fear memories in females.

Deconditioning-update group showed lower freezing expression throughout the reactivation sessions (Figure 2B). 24 hr after the last reactivation, both the footshock and no-footshock groups presented a comparable decrease in freezing behavior when compared to controls (Figure 2C). However, in the renewal and spontaneous recovery tests, fear expression reemerged in the no-footshock group, while the deconditioning-update group maintained its low freezing levels (Figure 2D and E, respectively; Supplementary file 2).

Next, we tested whether a stronger fear memory would constrain the deconditioning-update effect by training animals with 5 CS-US pairings using 1-mA shocks, while maintaining the rest of the protocol unchanged. In spite of the stronger shock intensity in the training session, the deconditioning-update group still presented reduced freezing levels in reactivation sessions when compared to the no-footshock group (Figure 2G). These lower freezing levels were maintained in the test, renewal and spontaneous recovery sessions, in which robust recovery was observed in no-footshock animals, but not in the deconditioning-update group (Figure 2H, I and J, respectively; Supplementary file 2). Similar results were observed in females in a slightly modified protocol with three reactivation sessions (Figure 2—figure supplement 1). These experiments suggest that the deconditioning-update induces a plastic state, allowing the fear memory trace to be altered even in boundary conditions that usually constrain memory updating (Pedraza et al., 2018).

In order to investigate whether the deconditioning-update approach is effective in attenuating other types of aversive memories, we trained animals in different fear-related tasks. First, we used a contextual fear conditioning paradigm, which is known to involve a set of brain regions that is partially distinct from auditory conditioning and includes the prefrontal cortex and hippocampus (Maren et al., 2013). Animals were placed in the conditioning chamber for 3 min before receiving two 0.5-mA, 2 s footshocks separated by a 30 s interval. On days 3 to 6, rats were reexposed to the same context for 4 min, with those in the deconditioning-update group receiving weak footshocks (0.1mA, 2 s) 3 min after being placed in the chamber (Figure 3A). The deconditioning-update group had lower fear expression during reactivations (Figure 3B) and maintained these lower freezing levels compared with the other groups in the test (Figure 3C). The same pattern was observed in the spontaneous recovery test, performed 20 days later, suggesting that the decrease in freezing caused by deconditioning-update is long-lasting (Figure 3D; Supplementary file 3).

Another set of animals underwent the step-through inhibitory avoidance task, in which training consists of applying 4 0.5-mA, 1-s footshocks with 5-s intervals when the animal enters the dark compartment of a shuttle box. During reactivations, animals were placed in the dark compartment for 30 s, with those in the deconditioning-update group receiving 2 0.1-mA shocks. In the test session, the animals were put in the light compartment, and the latency to enter the dark compartment was measured (Figure 3E). The deconditioning-update group showed a much shorter latency to reach the dark chamber (Figure 3F) and spent more time in the dark compartment over the 10-min session compared to the other groups (Figure 3G; Supplementary file 3), suggesting that memory was more robustly updated in this group during reactivation. Taken together, these results suggest that the deconditioning-update strategy is effective in weakening distinct types of fear-related memories.
Figure 3. Deconditioning-update weakens fear memory in different behavioral tasks. (A) Experimental design in contextual fear conditioning: rats were fear-conditioned with five contextual-shock pairings (4 min context + 4 US, 0.5mA). Starting 48 hr later, the no-footshock and footshock groups were exposed to daily reactivation sessions.

24 hr after the last reactivation, all groups were tested; 20 days later, they were tested for spontaneous recovery.

(B) Freezing levels during reactivation sessions. Rats exposed to weak footshocks during reactivation sessions showed a significant reduction in freezing responses maintained during the test (C) and spontaneous recovery (D) sessions. (E) Experimental design in inhibitory avoidance: rats were placed in the lighted compartment and received footshocks (4 US, 0.5mA) upon entering the dark one. Starting 48 hr later, the no-footshock and footshock groups were exposed to daily 30-s reactivation sessions in the dark compartment; 24 hr after the last reactivation, all groups were tested. Rats exposed to weak footshocks during reactivation sessions showed lower latencies to cross to the dark compartment (F) and spent more time in it during the test (G).

Bars represent mean ± SEM or median with interquartile range (in F and G). Statistical comparisons for contextual fear conditioning are performed using two-way repeated-measures ANOVA followed by a Bonferroni post-hoc (reactivation sessions) or one-way ANOVA followed by a Tukey post-hoc (test, renewal, and spontaneous recovery). For inhibitory avoidance, a Kruskal-Wallis test followed by a Dunn post-hoc was performed. *p<0.05; **p<0.005; ***p<0.0005; ****p<0.0001. For full statistics, see Supplementary file 3.

Figure 3 continued on next page
To address whether the deconditioning-update effect would also be observed within a single, long-lasting extinction session, we trained animals and exposed them 48 hr later to a session containing 12 CSs, with the deconditioning-update group receiving a 0.1-mA footshock at the end of each tone. Fear reduction was limited and largely similar across groups in the extinction session and in the subsequent tests (Figure 4—figure supplement 1), albeit with slightly lower renewal in the deconditioning-update group, suggesting that the pairing of the CS with a weak footshock is not as effective in accelerating single-session extinction. In order to further explore this possibility, we subjected animals to a 24-tone single-session extinction protocol (Figure 4A). This led to robust fear reduction both within the extinction session (Figure 4B) and in a subsequent test (Figure 4C) in the no-footshock group, while freezing remained largely unchanged in the footshock group. However, fear memory reemerged in the renewal and spontaneous recovery test among no-footshock animals (Figure 4D and E; Supplementary file 4), as typically occurs with extinction learning. These results show that the presence of a weak shock at the end of every CS not only does not enhance extinction occurring over a single behavioral session, but can actually impair it in the short-term.

Other studies have shown that activation of L-type voltage-gated Ca\(^{++}\) channels (LVGCC) is necessary both for destabilizing a reactivated memory during reconsolidation (Suzuki et al., 2008; Lee and Flavell, 2014; Crestani et al., 2015; Haubrich et al., 2015) and for some forms of extinction (Cain et al., 2002). Thus, we used the LVGCC antagonist nimodipine as a pharmacological tool to investigate whether deconditioning-update involved memory destabilization mechanisms (Flavell et al., 2011; Sierra et al., 2013; Crestani et al., 2015; Haubrich et al., 2015). Animals were trained and divided into four groups: no-footshock + vehicle, no-footshock + nimodipine, deconditioning-update + vehicle and deconditioning-update + nimodipine, with nimodipine or vehicle administered systemically 30 min before each reactivation (Figure 4F). Nimodipine attenuated freezing decrease in both behavioral protocols, suggesting that the drug impaired both deconditioning-update and regular extinction (Figure 4G and H). In the renewal and spontaneous recovery sessions, high freezing levels reemerged both in the no-footshock vehicle group and in nimodipine-treated animals, while the deconditioning-update vehicle group maintained a lower freezing level (Figure 4I and J; Supplementary file 4). This suggests that deconditioning-update is mediated by memory destabilization processes requiring Ca\(^{++}\) influx through LVGCCs.

In order to address whether the effect of nimodipine treatment before reactivation sessions might be explained by state-dependency of the extinction memory in the test session, we trained a new set of animals in the same condition, except that two tests were conducted on separate days in each animal, either in the presence or absence of nimodipine treatment 30 min before. We found that animals treated with nimodipine before the test kept expressing high freezing levels, suggesting that nimodipine injection before reactivations prevented memory destabilization instead of inducing state dependency (Figure 4—figure supplement 2).

One explanation for our findings is that pairing the CS with a weak footshock could lead to a smaller degree of prediction error during reexposure sessions, biasing them towards memory updating as opposed to new learning. Computational models using neural networks (Osan et al., 2011) or Bayesian approaches (Gershman et al., 2017) have explored how different degrees of mismatch between stored memories and new experiences can lead to these two outcomes, suggesting lower degrees of mismatch or prediction error could lead to greater destabilization of stored memories. With this in mind, we used an adaptation of one of these models (Osan et al., 2011) to explore if this framework could account for our main results – that is accelerated fear reduction over multiple sessions and lower memory recovery when mismatch is reduced during reexposure. This fully connected Hopfield-like network (Hopfield, 1984) of 100 neurons is capable of storing patterns using Hebbian learning rules and retrieving them according to the inputs presented, which in our simulations included neurons representing tone and context information, as well as shock/non-shock information (Figure 5A). Additionally, the network also updates synaptic weights according to mismatch between a cue input and the retrieved network pattern (Osan et al., 2011).
Figure 4. Deconditioning-update is based on memory destabilization mechanisms. (A) Experimental design: rats were fear-conditioned with five tone-shock pairings (context A; 5CS+US, 0.5mA). 48 hr later, the no-footshock and footshock groups underwent a single extinction session (context B, 24 CSs), followed by test (context B), renewal (context A) and spontaneous recovery (context B) sessions. (B) Freezing levels during extinction. Weak footshocks...
As shown in Figure 5B, a lower degree of mismatch in the reexposure pattern, caused by weak activation of shock-related neurons (as would be expected from deconditioning-update), can indeed lead to a greater reduction in subsequent fear behavior than a ‘pure extinction’ pattern (sharing only the tone information with the original memory). This effect was sensitive to blockade of the mismatch-induced degradation term – the model equivalent of blocking memory destabilization mechanisms such as LVGCCs with a drug like nimodipine (Figure 5C). This occurs because the ‘minor footshock’ pattern leads to recovery of the original memory as the outcome of the initial sessions (Figure 5D), causing mismatch-induced updating of the weights supporting these memories and weakening of the mutual connections between shock and context neurons (Figure 5E). On the other hand, no-footshock extinction leads to immediate formation of a new attractor, leaving the connection weights related to the shock memory unaltered throughout the extinction process.

Because of this, deconditioning-update was sensitive to blockade of mismatch-induced degradation, while no-footshock extinction – which depends basically on new learning – was not (Figure 5C). Blocking the Hebbian plasticity term, on the other hand, led to blockade of no-footshock extinction, as well as reconsolidation blockade in low-mismatch conditions (Figure 5—figure supplement 1). Interestingly, blocking Hebbian plasticity during deconditioning-update led to greater freezing decrease after the first reactivation session, suggesting that the procedure could be modulated by pharmacological agents such as protein synthesis inhibition; on the long run, however, the same manipulation decreased extinction due to interference with new learning.

A caveat here is that, unlike in the model, nimodipine did affect regular extinction in the experimental results. This suggests that the mechanistic distinction between deconditioning-update and classic extinction is not so clear-cut, and that these two processes might share common mechanisms, as has been proposed for reconsolidation and extinction (Almeida-Correão and Amaral, 2014). Capturing these subtleties, as well as other features of the data such as distinctions between single- and multiple-session extinction, likely requires model implementations that are more complex than this simple adaptation of the classic Hopfield formulation.

Discussion

Taken as a whole, our findings demonstrate that presenting a tone followed by a weak footshock (deconditioning-update) was more effective in reducing fear memory than presenting a tone in the absence of shock (extinction training). Attenuation of fear responses following deconditioning-update was robust, long-lasting, and less sensitive to renewal and spontaneous recovery. Remarkably, this strategy was also effective in reducing fear within boundary conditions in which memories have been described to be less sensitive to modification (e.g., very strong training protocols and remote memories). The same effect was found in other types of fear-related memory (contextual...
Figure 5. Lower mismatch accelerates fear reduction and decreases renewal in a neural network model. (A) Cue inputs presented to the network during training (shock memory), reexposure (with or without footshock) and test sessions (consisting of the tone and either context B (test) or A (renewal)). Color scale shows the cue received by each of the 100 neurons (B) Extinction over multiple sessions using the no-footshock (red bars) or footshock (blue bars) cue. Bars represent freezing, expressed as the activity ratio between shock neurons and the sum of shock and non-shock neurons in response to the test cue, at reexposure days 1 to 4. After each test, memory is updated according to the activity reached in response to the full reexposure pattern. (C) Effect of LGVCC blockade (i.e. setting the mismatch-induced degradation term D to 0). Removing the degradation term blocks deconditioning-update, but not regular extinction. (D) Network activity in retrieval tests during tone presentations (e.g. cued with the tone alone) and at the end of reexposure (e.g. cued with the full reactivation pattern), as well as on test and retrieval sessions. Lower mismatch (i.e. weak footshock) leads to retrieval of the original pattern on the first days, leading to memory updating through mismatch-induced degradation and lower retrieval on subsequent tests. (E) Mean synaptic weights between different neuronal groups after training and at the end of each extinction session. Heat map represents the connection from neuronal populations in the Y axis to those in the X axis in the no-footshock and footshock groups. Deconditioning-update leads to weakening of connections between context and shock neurons and of their inhibitory connections to other neurons. On no-footshock extinction, an extinction memory is formed with sparing of the shock representation.

The online version of this article includes the following figure supplement(s) for figure 5:

Figure supplement 1. Blockade of Hebbian plasticity impairs reconsolidation, blocks standard extinction and interacts with deconditioning-update.
fear conditioning and inhibitory avoidance tasks), and was sensitive to pharmacological blockade of LGVCCs. We suggest that the CS–US association is weakened during the deconditioning-update approach, leading to lower fear expression. Considering that (i) fear reduction is long-lasting, without spontaneous recovery, renewal and savings, (ii) weak-footshock pairings do not have the same effect in single-session extinction protocols or when stronger footshocks are applied, and (iii) this effect is prevented by LGVCC antagonism, we suggest that deconditioning-update is mediated by the memory destabilization effects commonly associated with reconsolidation.

These results are in line with predictions from a neural network model previously built to account for transitions between reconsolidation and extinction with increasing reexposure time in contextual fear conditioning (Osan et al., 2011). In this model, modification of existing connection weights is triggered by a protein synthesis-independent set of processes when there is mismatch between the memory pattern retrieved by the network (based on previously learned experiences) and the currently experienced sensory state, as when the prediction of a strong footshock is offset by a mild one. However, if mismatch is too extensive, as during extinction in the absence of a footshock, a new attractor is formed in the network at the first reexposure session already, preventing mismatch-induced weakening of the original memory.

Although the model is quite reductionist in its implementation, both due to the absence of realistic topology and to the non-spiking, continuous nature of neuronal activity, it is nevertheless able to account for transitions between simple retrieval, reconsolidation and extinction with increasing reexposure time (Suzuki et al., 2004) or number of non-reinforced CSs (Lee et al., 2006; Sevenster et al., 2014), as assessed by the effects of pharmacological agents such as protein synthesis inhibitors, NMDA antagonists and β-blockers. The model also helps to explain why, unlike pharmacological reconsolidation blockade, deconditioning-update takes several sessions to occur, as the impact of memory labilization mechanisms on a stored memory is much more pronounced when Hebbian mechanisms are blocked concomitantly (Osan et al., 2011).

Although we have not tested this explicitly, our results also appear compatible with the Bayesian inference framework proposed by Gershman et al. (2017), in which the probability that an experience is attributed to a new latent cause increases in proportion to the degree of prediction error generated by previous experience. This can explain why a lower degree of mismatch, such as that caused by reactivation followed by a weak footshock, can lead to a greater probability of memory updating and decrease the recovery of fear. If this is the case, the rationale for the greater effectiveness of deconditioning-update in reducing fear might be similar to that observed in so-called retrieval extinction paradigms (Monfils et al., 2009; Kredlow et al., 2016). In this case, the first retrieval trial, usually consisting of a single CS, is thought to induce retrieval-induced destabilization of the original memory, as the prediction error generated by this reexposure is not sufficient to form an extinction memory. Nevertheless, following this initial retrieval session with an extinction procedure leads to lower fear recovery than when extinction is performed without it, although this effect has been observed rather inconsistently across studies (Auber et al., 2013; Kredlow et al., 2016).

Interestingly, deconditioning-update only occurred when reactivation sessions were spaced across multiple days. When a massed extinction procedure with 24 non-reinforced CSs was performed within a single extinction session, on the other hand, a mild footshock at the end of the tones actually prevented within-session extinction, and increased freezing in a test performed on the following day. This seems to reinforce the notion that computations linking the degree of prediction error with memory destabilization occur only at the end of reexposure (Perez-Cuesta et al., 2007; Osan et al., 2011). It is also in line with the idea that within- and between-session extinction are distinct processes, with different dynamics and molecular requirements (Plendl and Woćjak, 2010; Almeida-Corrêa et al., 2015).

An important limitation of our computational framework, however, is that time is not explicitly modeled – thus, one cannot distinguish between massed and spaced extinction protocols in order to investigate the different results found in the two settings. An interesting challenge for future theoretical models, thus, would be to study whether and how within-session extinction relates mechanistically to mismatch-induced updating and between-session extinction. In this line, it is interesting to note that the model by Gershman et al. (2017) postulates that the probability that an experience is attributed to a new latent cause increases with time between initial learning and reactivation. This could plausibly account for why fear reduction with deconditioning-update is only observed in...
spaced sessions, in which new structural learning is more likely to take place in the absence of shocks (and thus prevent the original memory from updating in standard extinction conditions).

Nevertheless, in contrast to our work, Gershman et al. (2013) did find an effect of reducing mismatch at the start of a single 24-CS extinction session by pairing some of the initial tones with full-strength footshocks. That said, their effect was only observed in spontaneous recovery sessions and post-reinstatement sessions, and not within the extinction session itself. Moreover, the approach to induce lower degrees of mismatch in their experiment – gradually reducing the frequency of regular footshocks – was different from ours, in which this was achieved by providing a low-intensity shock at the end of every tone. Studying whether both approaches could be combined – by gradually decreasing footshock intensity, for example – could be an interesting topic for further investigation of the deconditioning-update effect. Another topic for future studies is whether mismatch between the training and reactivation context in auditory conditioning protocols also contributes to the deconditioning-update effect, as recently reported for reconsolidation (Ferrara et al., 2019b). In our study, animals showed relatively high pre-CS freezing levels, suggesting some degree of generalization between contexts, although the presence of renewal suggests that animals were also capable of differentiating them. It is thus an open question whether generalization might be important for deconditioning-update to occur.

Psychiatric disorders associated with pathological memories are prevalent, cause important social and economic burden, and approaches to translate basic knowledge of fear conditioning for potentiation of exposure therapy have met limited success so far in clinical settings (Monfils and Holmes, 2018). We believe that exploring the principles of deconditioning-update – that is, the notion that there is an ideal window of prediction error to potentiate reexposure effects – is a promising therapeutic avenue that could be explored in more depth in the setting of trauma-focused psychotherapy. The high efficacy, long-lasting effects, and drug-free nature of this approach make it particularly appealing for translation to human memory-related disorders, such as trauma, phobias and drug abuse.

Materials and methods

A total of 323 male and female Wistar rats (2–3 months old, weighing between 300 and 400 g) from CREAL at the Federal University of Rio Grande do Sul (UFRGS) were used for the experiments. Only one animal was excluded (in the experiment in Figure 1A) due to health conditions. They were housed in plexiglass boxes, with four animals per cage, with block randomization using the cage as subgroup to ensure that each cage contained at least one animal per experimental group. Sample sizes ranged from 6 to 10 per group across experiments, yielding statistical power estimates between 69% and 89% to detect an absolute difference of 30% in freezing time (in line with the differences in memory recovery observed in Monfils et al. (2009)) with a standard deviation of 20% (our average value for test sessions in Figure 1) at $\alpha = 0.05$ in a 2-tailed t test.

Animals were kept on 12/12 hr light/dark cycle under controlled temperature (21°C ± 2), with regular chow and water available ad libitum and humidity of approximately 65%. All experiments were performed during the light cycle. All procedures followed the Brazilian ethical guidelines for animal research set by the National Council for the Control of Experimental Animal Research (CONCEA). Methods and results are reported according to the revised ARRIVE guidelines (Percie du Sert et al., 2019).

Auditory fear conditioning

Apparatus: The conditioning chamber (context A) consisted of an illuminated plexiglass box (33 × 22 × 22 cm), with a floor grid of parallel 0.1-cm caliber stainless steel bars spaced 1 cm apart. All context chambers had the same dimensions, but context A had black walls, whereas context B was vertically striped in black and white. Context C consisted of white and brown lateral walls and a transparent front wall.

Training session: Rats were habituated for 2 days in context B (10 min each), and 24 hr later were placed in context A, where they received five conditioning trials consisting of a 30 s presentation of a 5-kHz, 75-dB tone (CS) that co-terminated with a 0.5-mA (or 1-mA in Figure 2F–H), 1-s footshock (US) (five tone-footshock pairings). The first CS was presented 2 min into the session, with an interpairing interval of 1 min. One minute after the final pairing, rats were returned to their home cages.
**Reactivation sessions**: In daily sessions taking place in context B and starting 48 hr after training (or 40 days later in Figure 2A–E), animals in the no-footshock group received 3 CS-only, while the footshock group (deconditioning-update) received 3 CSs that co-terminated with a 0.1-mA (or 0.3-mA in Figure 1—figure supplement 1), 1-s shock. The percentage of time freezing during each tone presentation was quantified and the mean freezing percentage for the three tones was used as a measure of fear. The first CS was presented 2 min into the session, with an interpairing interval of 1 min. One minute after the final pairing, rats were returned to their home cages. Most experiments used four reactivation sessions, except for those in Figure 1—figure supplement 2 (one session) and Figure 2—figure supplement 1 (three sessions). In the devaluation experiment (Figure 1—figure supplement 3A–B), the protocol was the same, except that the 0.1-mA footshocks were presented in context C without being paired with the tone. In the paired vs. unpaired experiment (Figure 1—figure supplement 4), the unpaired group received uncorrelated tones and 0.1-mA footshock in a pseudorandom order. Pre-CS freezing levels were measured for the 30 s immediately preceding the first tone.

**Test session**: 24 hr after the last reactivation session, both groups were presented with 3 CSs in context B. The percentage of time freezing during each tone presentation was quantified, and the average for the three tones was used as a fear measure. The first CS was presented 2 min into the session, with an interpairing interval of 1 min. One minute after the final pairing, rats were returned to their home cages. Pre-CS freezing levels were measured for the 30 s immediately preceding the first tone.

**Renewal**: 24 hr after the test session, animals were placed in context A, where they received 3 CSs. The percentage of time freezing during each tone presentation was quantified, and the average percentage was used as a measure of fear recovery. The first CS was presented 2 min into the session, with an interpairing interval of 1 min. One minute after the final pairing, rats were returned to their home cages. Pre-CS freezing levels were measured for the 30 s immediately preceding the first tone.

**Spontaneous Recovery**: 20 days after the renewal session, animals were placed in context B and received 3 CSs. The percentage of time freezing during each tone presentation was quantified, and the average was used as a measure of fear recovery. The first CS was presented 2 min into the session, with an interpairing interval of 1 min. One minute after the final pairing, rats were returned to their home cages. Pre-CS freezing levels were measured for the 30 s immediately preceding the first tone.

**Reinstatement**: 24 hr after the test session, animals were exposed to context C for 5 s, where they received two 2-s, 0.7-mA footshocks. 24 hr later, they were tested for reinstatement in context B. Pre-CS freezing levels were measured for the 30 s immediately preceding the first tone.

**Retraining**: 24 hr after the spontaneous recovery test, rats were submitted to a training procedure (3 × 0.5 mA). They were then tested 24 hr later to assess savings. Pre-CS freezing levels were measured for the 30 s immediately preceding the first tone.

**Single-session extinction**: Animals were placed in context B 48 hr after training, where they received either 12 or 24 CSs depending on the protocol. In the no-footshock group these tones were not accompanied by the US, while the footshock group received tones that co-terminated with a 0.1-mA footshock. The first CS was presented 2 min into the session, with an interpairing interval of 1 min. One minute after the final pairing, rats were returned to their home cages. Pre-CS freezing levels were measured for the 30 s immediately preceding the first tone.

**Contextual fear conditioning**

**Apparatus**: The conditioning chamber consisted of an illuminated plexiglass box (33 × 22 × 22 cm grid of parallel 0.1-cm caliber stainless steel bars spaced 1 cm apart).

**Training session**: In the training session, rats were placed in the conditioning chamber for 3 min before receiving two 2-s, 0.5-mA footshocks separated by a 30 s interval; they were kept in the conditioning context for an additional 30 s before returning to their home cage.

**Reactivation session**: 48 hr after the training session, animals were reexposed to the same conditioning chamber for 4 daily 4 min sessions. Rats from the footshock group received two pairs of 0.1-mA, 2-s shocks after 180 and 210 s, while the no-footshock group did not receive any shocks.
Test session and spontaneous recovery: 24 hr after the last reactivation session, animals were re-exposed to the same conditioning chamber for a 4 min test session and the percentage of time freezing was quantified. 20 days later, the procedure was repeated to assess spontaneous recovery.

Step-through inhibitory avoidance

**Apparatus:** The apparatus consists of an automated box (Insight Ltda., Brazil) with two compartments, a dark one and a lighted one, each measuring $33 \times 22 \times 22$ cm. The floor consisted of a grid of metal bars with 1 mm diameter placed 1 cm from each other.

**Training session:** Animals were placed in the lighted compartment. When they entered the dark compartment, the door was closed and the animals received 4 0.5-mA, 1-s footshocks, with intervals of 5 s between them. They were removed from the box 10 s after the last footshock.

**Reactivation sessions:** In daily sessions starting 48 hr after training, animals in the footshock and no-footshock groups were placed in the dark compartment for 30 s, with no access to the lighted compartment. The footshock group received 2 0.1-mA shocks (at 25 s and 30 s) while the no-footshock group did not receive any shocks.

**Test session:** All animals were placed in the lighted compartment and left free to explore the box. The latency to enter into the dark compartment for the first time and the time spent in each compartment were counted over a 10 min session and used as measures of memory.

Behavioral assessment

Freezing behavior was used as a memory index in the fear conditioning tasks, being registered with a stopwatch in real time by an experienced observer that was blinded to the experimental group. Freezing was defined as total cessation of all movements except those required for respiration.

Open field

Exploratory activity and anxiety-like behavior were assessed in the open field test in order to exclude non-specific effects of nimodipine. The apparatus consisted of a circular arena (90-cm diameter) with 50-cm high walls. The floor was subdivided into 12 quadrants and three concentric zones (periphery, intermediary and center). Animals were exposed to the apparatus for 5 min, during which the time spent on the periphery (thigmotaxis) and the number of crossings between quadrants were measured. Nimodipine (16 mg/kg) or vehicle was measured intraperitonally 30 min before the test.

Drugs

Nimodipine (Sigma), an antagonist of the L-type voltage-gated calcium channels (LGVCCs) was dissolved in sterile isotonic saline solution with 8% dimethylsulfoxide to a concentration of 16 mg/mL. Nimodipine or its vehicle was injected intraperitoneally at a volume of 1 mL/kg (16 mg/kg) 30 min before memory reactivation sessions, test sessions or open field sessions.

Statistical analysis

Data are expressed as mean ± SEM, always using the animal as the experimental unit. The statistical tests used and their results are detailed for every experiment in Supplementary file 1–11 and in Figure 1—source data 1, Figure 2—source data 1, Figure 3—source data 1 and Figure 4—source data 1; they include two-tailed Student’s t test; one-way, two-way or three-way analysis of variance (ANOVA), followed by Tukey’s or Bonferroni’s post hoc test, when necessary; and Kruskal-Wallis test, followed by Dunn’s post hoc. Values of $p<0.05$ were considered statistically significant. Baseline freezing levels for all experiments are shown on Supplementary file 12–21. Unit-level data for all figures are provided as Figure 1—source data 1, Figure 2—source data 1, Figure 3—source data 1 and Figure 4—source data 1.

Computational simulations

Model Network: In order to propose a mechanistic explanation for the experimental results, we used an adaptation of the attractor network model described in Osan et al. (2011). This Hopfield-like network is capable of storing and retrieving memories using Hebbian learning rules dependent on neuronal activity, which in turn depends on the inputs presented to a fully connected network of 100 neurons. In this network, the activity of each neuron $i$ is determined by
\[
\tau \frac{du_i}{dt} = -u_i + \frac{1}{2} \left( 1 + \tanh \left( \sum_{j=1}^{N} w_{ij} u_j + I_i \right) \right)
\]

where \( \tau \) is the neural time constant and \( u_i \) represents the level of activation of neuron \( i \) which can vary continuously from 0 to 1 – unlike in the original Hopfield continuous activity model \((\text{Hopfield}, 1984)\), in which activity varies from \(-1\) to 1.

As a fully connected neural network, every neuron \( i \) is connected with every neuron \( j \). For the learning process, the network needs to reinforce the connections between neurons that fire together, while creating inhibition when presynaptic neuron \( i \) is active and postsynaptic \( j \) is silent. Changes in the synaptic weight matrix \( W = (w_{ij}) \) are determined by the equation

\[
\Delta W = -\gamma W + \text{HLP} + \text{MID}
\]

where Hebbian learning plasticity (HLP) and mismatch-induced degradation are two independent learning rules (see below) and \( 0 \leq \gamma \leq 1 \) is a time-dependent synaptic decay factor.

Learning occurs by presenting an input \( I_i \) to the circuit, corresponding to sensory information provided by the environment and/or internal cues, which lead to changes in the plastic connections between neurons. The cue has a one-to-one topology to the memory network, with every neuron receiving a cue input that can be either excitatory or inhibitory. Modifications on the synaptic weight matrix follow the HLP rule, corresponding to the Hebbian formulation implemented in classic Hopfield networks and described as

\[
\text{HLP} = S(u^T + u) - S((1 - u)^T + u)
\]

where vector \( u = (u_1, u_2, \ldots, u_N) \) is the steady state of the network after input \( I_i \) presentation, while \( S \) is a factor representing requirements for Hebbian plasticity, such as protein synthesis, receptor activation, intracellular signaling and other mechanisms.

When the cue input leads to the retrieval of a previously stored memory, this can cause mismatch between the cue input and the retrieved attractor if the two are not the same. This leads to concomitant activation of the MID learning rule, corresponding to a memory-updating system akin to that involved in memory destabilization during reconsolidation and defined by

\[
\text{MID} = D(m^T + u)
\]

where \( D \) is a factor representing requirements for memory destabilization (such as protein degradation and LVGCCs), \( m = I_{\text{norm}} - u \) is the mismatch vector defined and \( I_{\text{norm}} \) is a normalized cue vector. The MID term leads to weakening of connections responsible for the mismatch in order to update the existing memory.

**Learning, retrieval and reactivation**

Non-overlapping neuron clusters in the network were chosen to represent the training or extinction contexts (six neurons each), tone (two neurons), aversive stimulus/shock (10 neurons) or safety/absence of shock (10 neurons) (Figure 5A). Initially, a pattern representation of a memory unrelated to fear conditioning was presented as a cue to the network. This was followed by a training pattern activating neurons representing context A, tone and shock while inhibiting the remaining ones.

Retrieval was evaluated at every training or reactivation session through presentation of a cue activating neurons representing context B and tone, with no input to the remaining neurons. This corresponds to the period in which freezing is assessed (e.g. during the tone itself, shown as ‘tone’ in Figure 5D), and was modeled with the same cue irrespectively of the presence of shock at the end of reactivation. For the renewal test, the retrieval cue activated neurons representing context A and tone. To quantify memory retrieval, we used the mean activity of neurons representing shock and absence of shock, which was converted to a ‘freezing percentage’ by dividing the total activity of shock neurons by the total activity of both groups – thus, 100% freezing corresponds to full activation of shock neurons and no activation of non-shock neurons.

At the end of each reactivation session, the network underwent a new learning round with a pattern that varied according to the experimental group. To model standard extinction over multiple retrieval sessions, we activated the non-shock neuron cluster along with the neurons representing
the extinction context and tone, while inhibiting the remaining ones (‘no-footshock reactivation’ in Figure 5A). For deconditioning-update, we assumed an intermediate representation between the learning and extinction patterns (‘footshock reactivation’ in Figure 5A). For reconsolidation (Figure 5—figure supplement 1), this intermediate representation was closer to the shock pattern than to the extinction one. Synaptic weights were updated according to the activation pattern reached in response to these cues (shown as ‘session end’ in Figure 5D). Unlike in the original model, no synaptic decay was assumed (i.e. \( \gamma \) was set to 0) and learning strength (as defined by \( S \)) was assumed to be smaller during reactivation sessions in both groups due to the lower intensity of the stimuli, thus allowing extinction to occur over multiple sessions.

After each learning or reactivation session, the mean synaptic weight between each cluster of neurons (tone, contexts, shock and non-shock) was calculated by taking the average of the connections between all presynaptic neurons of a subpopulation and all postsynaptic neurons of the other subpopulation. This was used to create the synaptic weight matrix between clusters shown in Figure 5E.

**Model parameters**

All simulations were performed in MATLAB R2018a (Mathworks) using \( N = 100; \tau = 1; \gamma = 0; s_0 = 1 \). For training sessions, we set \( S = 0.8 \), while in reactivation sessions we used \( S = 0.25 \). \( D \) was set to 0.95 for all sessions, except for reactivations using nimodipine, in which \( D = 0 \). Each unit \( i \) during learning received an input \( I_i \) varying between \(-5\) and 5. In the deconditioning update group, the aversive shock cluster received an input \( I_i \) of \(-2.31\), while non-shock neurons received 2.31 [corresponding to \( t = 6 \) in the transformation used by Osan et al. (2011) to create intermediate patterns]. For reconsolidation, these inputs were 3.80 and \(-3.80\), respectively, corresponding to \( t = 3 \) in Osan et al. (2011). In the retrieval cue, each targeted neuron had an input \( I_j = 1.5 \) and 0 for other neurons.

One hundred simulations of each experiment were performed, with different initial conditions determined by Gaussian noise in the initial weight matrices (with a normal distribution on \([-0.05, 0.05]\)) and in the neuronal activation at the start of every session (with a normal distribution on \([0, 0.1]\)). For each simulation, 100 retrieval trials were run in each session to determine freezing percentage. All results are displayed as the mean ± S.E.M of these 100 simulations. Matlab code to perform all simulations and generate Figure 5B, C and E and Figure 5—figure supplement 1 is presented as Source Code.

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Ethics
Animal experimentation: All procedures followed the Brazilian ethical guidelines for animal research set by the National Council for the Control of Experimental Animal Research (CONCEA) and approved by the committee on the Ethics of Animal Experiments of the UFRGS (number 34547).

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Additional files

Supplementary files
- Source code 1. Matlab code for Figure 5.
- Supplementary file 1. Weakening fear memory through deconditioning-update training.
- Supplementary file 2. Deconditioning-update approach weakens both remote and strong fear memory.
- Supplementary file 3. Deconditioning-updating weakens fear memory in different behavioral tasks.
- Supplementary file 4. Deconditioning-update is based on memory destabilization mechanisms.
- Supplementary file 5. Deconditioning-update does not occur with 0.3-mA shocks.
- Supplementary file 6. A single reactivation session does not update fear memory.
- Supplementary file 7. Deconditioning-update is not due to US devaluation.
- Supplementary file 8. Deconditioning-update does not occur with unpaired shocks in the reactivation sessions.
- Supplementary file 9. Deconditioning-update weakens strong fear memories in females.
- Supplementary file 10. Deconditioning-update does not occur in a single 12-CS extinction session.
- Supplementary file 11. Nimodipine does not affect open field behavior and does not induce a state-dependent memory.
- Supplementary file 12. Baseline (pre-CS) freezing levels for Figure 1.
- Supplementary file 13. Baseline (pre-CS freezing levels for Figure 2.
- Supplementary file 14. Baseline (pre-CS) freezing levels for Figure 4.
- Supplementary file 15. Baseline (pre-CS) freezing levels for Figure 1—figure supplement 1.
- Supplementary file 16. Baseline (pre-CS) freezing levels for Figure 1—figure supplement 2.
- Supplementary file 17. Baseline (pre-CS) freezing levels for Figure 1—figure supplement 3.
- Supplementary file 18. Baseline (pre-CS) freezing levels for Figure 1—figure supplement 4.
- Supplementary file 19. Baseline (pre-CS) freezing levels for Figure 2—figure supplement 1.
- Supplementary file 20. Baseline (pre-CS) freezing levels for Figure 4—figure supplement 1.
- Supplementary file 21. Baseline (pre-CS) freezing levels for Figure 4—figure supplement 2.
Data availability

All data generated or analysed during this study are included in the manuscript and supporting files.

References

Almeida-Corrêa S, Moulin TC, Carneiro CF, Gonçalves MM, Junqueira LS, Amaral OB. 2015. Calcineurin inhibition blocks within- but not between-session fear extinction in mice. Learning & Memory 22:159–169. DOI: https://doi.org/10.1101/lm.037770.114, PMID: 25691516

Almeida-Corrêa S, Amaral OB. 2014. Memory labilization in reconsolidation and extinction—evidence for a common plasticity system? Journal of Physiology-Paris 108:292–306. DOI: https://doi.org/10.1016/j.jphysparis.2014.08.006, PMID: 25173958

Archbold GE, Bouton ME, Nader K. 2010. Evidence for the persistence of contextual fear memories following immediate extinction. European Journal of Neuroscience 31:1303–1311. DOI: https://doi.org/10.1111/j.1460-9568.2010.07161.x, PMID: 20345921

Auber A, Tedesco V, Jones CE, Monfils MH, Chaimulera C. 2013. Post-retrieval extinction as reconsolidation interference: methodological issues or boundary conditions? Psychopharmacology 226:631–647. DOI: https://doi.org/10.1007/s00213-013-3004-1, PMID: 23404065

Beckers T, Kindt M. 2017. Memory reconsolidation interference as an emerging treatment for emotional disorders: strengths, limitations, challenges, and opportunities. Annual Review of Clinical Psychology 13:99–121. DOI: https://doi.org/10.1146/annurev-clinpsy-028161-045209, PMID: 28375725

Bouton ME. 2002. Context, ambiguity, and unlearning: sources of relapse after behavioral extinction. Biological Psychiatry 52:976–986. DOI: https://doi.org/10.1016/S0006-3223(02)01546-9, PMID: 12437938

Bouton ME, Winterbauer NE, Todd TP. 2012. Relapse processes after the extinction of instrumental learning: renewal, resurgence, and reacquisition. Behavioural Processes 90:130–141. DOI: https://doi.org/10.1016/j.beproc.2012.03.004, PMID: 22450305

Bustos SG, Maldonado H, Molina VA. 2009. Disruptive effect of midazolam on fear memory reconsolidation: decisive influence of reactivation state and memory age. Neuropsychopharmacology 34:446–457. DOI: https://doi.org/10.1038/npp.2008.75, PMID: 18509330

Cain CK, Blouin AM, Barad M. 2002. L-type voltage-gated calcium channels are required for extinction, but not for acquisition or expression, of conditional fear in mice. The Journal of Neuroscience 22:9113–9121. DOI: https://doi.org/10.1523/JNEUROSCI.22-20-09113.2002, PMID: 12388619

Crestani AP, Zacouteguy Boos F, Haubrich J, Ordóñez Sierra R, Santana F, Molina JM, Cassini LF, Alves LO, Quillfeldt JA. 2015. Memory reconsolidation may be disrupted by a distractor stimulus presented during reactivation. Scientific Reports 5:13633. DOI: https://doi.org/10.1038/srep13633, PMID: 26328547

De Oliveira Alves L, Crestani AP, Cassini LF, Haubrich J, Santana F, Quillfeldt JA. 2013. Reactivation enables memory updating, precision-keeping and strengthening: exploring the possible biological roles of reconsolidation. Neuroscience 244:42–48. DOI: https://doi.org/10.1016/j.neuroscience.2013.04.005, PMID: 23587841

Duvarci S, Nader K. 2004. Characterization of fear memory reconsolidation. Journal of Neuroscience 24:9269–9275. DOI: https://doi.org/10.1523/JNEUROSCI.2971-04.2004, PMID: 15496662

Fernández RS, Bavassi L, Kaczor L, Forcato C, Pedreira ME. 2016. Interference conditions of the reconsolidation process in humans: the role of Valence and different memory systems. Frontiers in Human Neuroscience 10:641. DOI: https://doi.org/10.3389/fnhum.2016.00641, PMID: 28066212

Ferrara NC, Jerome TJ, Cullen PK, Orsi SA, Kwapis JL, Trask S, Pullins SE, Helmstetter FJ. 2019a. GluR2 endocytosis-dependent protein degradation in the amygdala mediates memory updating. Scientific Reports 9:5180. DOI: https://doi.org/10.1038/s41598-019-41526-1, PMID: 30914678

Ferrara NC, Trask S, Pullins SE, Helmstetter FJ. 2019b. The dorsal Hippocampus mediates synaptic destabilization and memory lability in the amygdala in the absence of contextual novelty. Neurobiology of Learning and Memory 166:107089. DOI: https://doi.org/10.1016/j.nlm.2019.107089, PMID: 31563610

Flavell CR, Barber DJ, Lee JL. 2011. Behavioural memory reconsolidation of food and fear memories. Nature Communications 2:504. DOI: https://doi.org/10.1038/ncomms1515, PMID: 22009036

Frankland PW, Ding HK, Takahashi E, Suzuki A, Kida S, Silva AJ. 2006. Stability of recent and remote contextual fear memory. Learning & Memory 13:451–457. DOI: https://doi.org/10.1101/lm.183406

Gershman SJ, Jones CE, Norman KA, Monfils MH, Niv Y. 2013. Gradual extinction prevents the return of fear: implications for the discovery of state. Frontiers in Behavioral Neuroscience 7:164. DOI: https://doi.org/10.3389/fnbeh.2013.00164, PMID: 24302899

Gershman SJ, Monfils MH, Norman KA, Niv Y. 2017. The computational nature of memory modification. eLife 6:e23763. DOI: https://doi.org/10.7554/eLife.23763, PMID: 28294944

Haubrich J, Crestani AP, Cassini LF, Santana F, Sierra RO, Alves LO, Quillfeldt JA. 2015. Reconsolidation allows memory to be updated to a less aversive level through the incorporation of appetitive information. Neuropsychopharmacology 40:315–326. DOI: https://doi.org/10.1038/npp.2014.174, PMID: 25027331

Hopfield JJ. 1984. Neurons with graded response have collective computational properties like those of two-state neurons. PNAS 81:3088–3092. DOI: https://doi.org/10.1073/pnas.81.10.3088, PMID: 6587342
Kredlow MA, Unger LD, Otto MW. 2016. Harnessing reconsolidation to weaken fear and appetitive memories: a meta-analysis of post-retrieval extinction effects. Psychological Bulletin 142:314–336. DOI: https://doi.org/10.1037/bul0000034, PMID: 26689086

Lee JL, Milton AL, Everitt BJ. 2006. Reconsolidation and extinction of conditioned fear: inhibition and potentiation. The Journal of Neuroscience 26:10051–10056. DOI: https://doi.org/10.1523/JNEUROSCI.2466-06.2006, PMID: 17005868

Lee JL. 2009. Reconsolidation: maintaining memory relevance. Trends in Neurosciences 32:413–420. DOI: https://doi.org/10.1016/j.tins.2009.05.002, PMID: 19640595

Lee JL, Flavell CR. 2014. Inhibition and enhancement of contextual fear memory destabilization. Frontiers in Behavioral Neuroscience 8:144. DOI: https://doi.org/10.3389/fnbeh.2014.00144, PMID: 24808841

Maren S, Phan KL, Liberonz I. 2013. The contextual brain: implications for fear conditioning, extinction and psychopathology. Nature Reviews Neuroscience 14:417–428. DOI: https://doi.org/10.1038/nrn3492, PMID: 23635870

Milekic MH, Alberini CM. 2002. Temporally graded requirement for protein synthesis following memory reactivation. Neuron 36:521–525. DOI: https://doi.org/10.1016/S0896-6273(02)00270-0, PMID: 12008853

Monfils MH, Cowansage KK, Klann E, LeDoux JE. 2009. Extinction-reconsolidation boundaries: key to persistent attenuation of fear memories. Science 324:951–955. DOI: https://doi.org/10.1126/science.1167975, PMID: 19342552

Monfils MH, Holmes EA. 2018. Memory boundaries: opening a window inspired by reconsolidation to treat anxiety, trauma-related, and addiction disorders. The Lancet Psychiatry 5:1032–1042. DOI: https://doi.org/10.1016/S2215-0366(18)30270-0, PMID: 30385214

Nader K, Schafe GE, Le Doux JE. 2000. Fear memories require protein synthesis in the amygdala for reconsolidation after retrieval. Nature 406:722–726. DOI: https://doi.org/10.1038/35021052, PMID: 10963596

Nader K, Einassar EO. 2010. Memory reconsolidation: an update. Annals of the New York Academy of Sciences 1191:27–41. DOI: https://doi.org/10.1111/j.1749-6632.2010.05443.x, PMID: 20392274

Osan R, Tort AB, Amaral OB. 2011. A mismatch-based model for memory reconsolidation and extinction in attractor networks. PLOS ONE 6:e23113. DOI: https://doi.org/10.1371/journal.pone.0023113, PMID: 21826231

Pedraza LK, Sierra RO, Lotz FN, de Oliveira Alvares L. 2018. Periodical reactivation under the effect of caffeine attenuates fear memory expression in rats. Scientific Reports 8:7260. DOI: https://doi.org/10.1038/s41598-018-25648-6, PMID: 29740084

Percie du Sert N, Hurst V, Aihuvalia A. 2019. The ARRIVE guidelines 2019: updated guidelines for reporting animal research. bioRxiv. DOI: https://doi.org/10.1101/703181

Perez-Cuesta LM, Hepp Y, Pedreira ME, Maldonado H. 2007. Memory is not extinguished along with CS presentation but within a few seconds after CS-offset. Learning & Memory 14:101–108. DOI: https://doi.org/10.1101/lm.413507

Plendl W, Wojtak JT. 2010. Dissociation of within- and between-session extinction of conditioned fear. Journal of Neuroscience 30:4990–4998. DOI: https://doi.org/10.1523/JNEUROSCI.6038-09.2010, PMID: 20371819

Rescorla RA. 1973. Effect of US habituation following conditioning. Journal of Comparative and Physiological Psychology 82:137–143. DOI: https://doi.org/10.1037/h0033815, PMID: 4684968

Rescorla RA, Heth CD. 1975. Reinstatement of fear to an extinguished conditioned stimulus. Journal of Experimental Psychology: Animal Behavior Processes 1:88–96. DOI: https://doi.org/10.1037/0097-7403.1.1.88

Sevenster D, Beckers T, Kindt M. 2014. Prediction error demarcates the transition from retrieval, to reconsolidation, to new learning. Learning & Memory 21:580–584. DOI: https://doi.org/10.1101/lm.035493.114, PMID: 25320349

Sierra RO, Cassini LF, Santana F, Crestani AP, Duran JM, Haubrich J, de Oliveira Alvares L, Quillfeldt JA. 2013. Reconsolidation may incorporate state-dependency into previously consolidated memories. Learning & Memory 20:379–387. DOI: https://doi.org/10.1101/lm.030233.112, PMID: 23782508

Suzuki A, Josselyn SA, Frankland PW, Masushige S, Silva AJ, Kida S. 2004. Memory reconsolidation and extinction have distinct temporal and biochemical signatures. Journal of Neuroscience 24:4787–4795. DOI: https://doi.org/10.1523/JNEUROSCI.5491-03.2004, PMID: 15152039

Suzuki A, Mukawa T, Tsukagoshi A, Frankland PW, Kida S. 2008. Activation of LgVGCs and CB1 receptors required for destabilization of reactivated contextual fear memories. Learning & Memory 15:426–433. DOI: https://doi.org/10.1101/lm.035493.114, PMID: 18511694

Wang SH, de Oliveira Alvares L, Nader K. 2009. Cellular and systems mechanisms of memory strength as a constraint on auditory fear reconsolidation. Nature Neuroscience 12:905–912. DOI: https://doi.org/10.1038/nn.2350, PMID: 19543280