Extinction-Reconsolidation Boundaries: Key to Persistent Attenuation of Fear Memories
Marie-H. Monfils, et al.
Science 324, 951 (2009);
DOI: 10.1126/science.1167975

The following resources related to this article are available online at www.sciencemag.org (this information is current as of May 15, 2009):

Updated information and services, including high-resolution figures, can be found in the online version of this article at:
http://www.sciencemag.org/cgi/content/full/324/5929/951

Supporting Online Material can be found at:
http://www.sciencemag.org/cgi/content/full/1167975/DC1

This article cites 35 articles, 15 of which can be accessed for free:
http://www.sciencemag.org/cgi/content/full/324/5929/951#otherarticles

This article appears in the following subject collections:
Psychology
http://www.sciencemag.org/cgi/collection/psychology

Information about obtaining reprints of this article or about obtaining permission to reproduce this article in whole or in part can be found at:
http://www.sciencemag.org/about/permissions.dtl
Extinction-Reconsolidation Boundaries: Key to Persistent Attenuation of Fear Memories

Marie-H. Monfils,1,2,4 Kirilana K. Cowanage,1 Eric Klann,3 Joseph E. LeDoux1,3,4,5

Dysregulation of the fear system is at the core of many psychiatric disorders. Much progress has been made in uncovering the neural basis of fear learning through studies in which associative emotional memories are formed by pairing an initially neutral stimulus (conditioned stimulus, CS; e.g., a tone) to an unconditioned stimulus (US; e.g., a shock). Despite recent advances, the question of how to persistently weaken aversive CS-US associations, or dampen traumatic memories in pathological cases, remains a major dilemma. Two paradigms (blockade of reconsolidation and extinction) have been used in the laboratory to reduce acquired fear. Unfortunately, their clinical efficacy is limited: Reconsolidation blockade typically requires potentially toxic drugs, and extinction is not permanent. Here, we describe a behavioral design in which a fear memory in rats is destabilized and reinterpreted as safe by presenting an isolated retrieval trial before an extinction session. This procedure permanently attenuates the fear memory without the use of drugs.

When fearful memories are formed, they are initially labile but become progressively consolidated into persistent traces via the synthesis of new proteins (1, 2). Later retrieval of a consolidated fear memory engages two seemingly opposing mechanisms: reconsolidation and extinction (3–6). In the process of reconsolidation, a retrieved memory transiently returns to a labile state and requires new protein synthesis to persist further. During this labile state, the memory is amenable to enhancement or disruption (4, 7). The period of instability or lability, the reconsolidation window, persists for several hours after retrieval (8). Reconsolidation occurs in a broad range of learning paradigms (aversive and appetitive conditioning, explicit and implicit memory) (5, 9) and species (from snails to humans) (10, 11). Its adaptive purpose might be to enable the integration of new information present at the time of retrieval into an updated memory representation (4, 12, 13).

The possibility that reactivated memories may be modifiable was proposed many years ago (14, 15), and since then, numerous studies have demonstrated that blockade of the updating process engaged during retrieval—usually via pharmacological intervention within the reconsolidation window—prevents memory restoration and produces amnesia (loss of the specific memory that was reactivated in the presence of the drug or access to it) (4, 8, 12, 13). Thus, in the case of aversive memories, blocking reconsolidation weakens the emotional impact of a once fear-inducing stimulus by altering the molecular composition of the memory trace. This process generally requires the use of drugs that often cannot be readily administered to humans.

In contrast, fear extinction—a paradigm in which the conditioned stimulus (CS) is repeatedly presented in the absence of the unconditioned stimulus (US)—leads to progressive reduction in the expression of fear, but is not permanent because extinction does not directly modify the existing memory but instead leads to the formation of a new memory that suppresses activation of the initial trace (16–22). The efficacy of this inhibition, however, is strongly contingent on spatial, sensory, and temporal variables. Specifically, the reemergence of a previously extinguished fear is known to occur, in rodents and humans alike, under three general conditions: (i) renewal, when the CS is presented outside of the extinction context (17, 18); (ii) reinstatement, when the original US is given unexpectedly (19–22); or (iii) spontaneous recovery, when a substantial amount of time has passed (16, 17, 23). In clinical settings, where extinction-based exposure therapy is widely used as treatment for a number of anxiety-related disorders, including phobias and post-traumatic stress, exposure treatments are effective in some cases [e.g., (24, 25)]; however, they do not benefit everyone, and of those who do benefit, many show a return of fear due to spontaneous recovery, reinstatement, or renewal (18, 23, 26, 27).

Here, we devised an effective, drug-free paradigm for the persistent reduction of learned fear, capitalizing on differences between reconsolidation and extinction. Given that extinction training reduces the threatening value of the CS, we reasoned that when applied within the reconsolidation window (after the memory is rendered unstable by presenting an isolated retrieval trial), extinction training would result in the storage of the new nonthreatening meaning of the CS and prevent renewal, reinstatement, and spontaneous recovery, thus resulting in a more enduring reduction in fear relative to extinction training conducted outside the reconsolidation window. Specifically, we predicted that an extinction session presented after an isolated retrieval trial would lead to a persistent revaluation of the CS as less threatening, and/or a weakening of the stored trace or access to it, and thus would prevent the return of fear in the three aforementioned tests.

Six experiments were conducted. We first examined whether our behavioral paradigm could prevent the return of fear on a spontaneous recovery test, and if so, whether the observed effect was the result of an update during reconsolidation. We specifically designed this experiment on the basis of the premise that the lability window engaged at the time of retrieval is temporary—in rat fear conditioning, it closes within 6 hours (9)—at which time the memory is thought to be reconsolidated (4). We posited that if the interval between the isolated retrieval cue and extinction training was brief enough to enable the repeated unreinforced CSs to be presented within the lability window, then the new interpretation of the CS as no longer threatening should be incorporated during reconsolidation. If, however, the interval between the isolated retrieval trial and the beginning of extinction was outside the lability boundary, standard extinction should take place (i.e., rather than targeting the initial fear memory during its reconsolidation, a new memory would be formed in parallel with it and would act to temporarily suppress it) and fear should reemerge.

Rats were fear-conditioned using three tone-shock pairings, and were then divided into five experimental groups. Two groups had a retrieval-extinction interval within the reconsolidation window (10 min (n = 8) and 1 hour (n = 8)) and two groups outside the reconsolidation window (6 hours (n = 8) and 24 hours (n = 8)). In addition to these four retrieval (Ret) groups, a fifth group (No Ret) was exposed to context but did not receive a CS retrieval (n = 12). All procedures were conducted in context A (grid floor). All groups showed equivalent freezing for the last four trials of extinction [between-subjects analysis of variance (ANOVA), P > 0.1; Fig. 1]. Twenty-four hours later, all groups received a long-term memory (LTM) test to assess consolidation of extinction; the groups did not differ from one another (repeated-measures ANOVA, P > 0.1). All groups were tested 1 month after extinction, and their freezing to the CS was compared to their respective freezing at the 24-hour time point. A repeated-measures ANOVA revealed a Group × Time interaction, which suggested that there was a differential effect between the groups during the 24-hour LTM test and the 1-month test [F(1,39) = 22.47, P < 0.0001]. Simple main effects were then conducted to look at each group individually. The Ret groups with a retrieval-extinction interval outside the reconsolidation window, as well as the No Ret group, showed increased freezing (spontaneous recovery) relative to the 24-hour LTM test (within-subjects,
two-tailed t tests; no retrieval, t(11) = 5.225, P < 0.0001; 6 hours, t(11) = 5.671, P = 0.001; 24 hours, t(11) = 2.681, P = 0.031]; however, the Ret groups with an interval within the lability window did not [within-subjects, two-tailed t tests; 10 min, t(7) = 0.146, P = 0.888; 1 hour, t(7) = 1.59, P = 0.156] (28). These data are consistent with an update during reconsolidation.

To more fully address whether our procedure could prevent the return of fear, we further examined its effect on two additional assays: renewal and reinstatement. Two groups of rats were fear-conditioned as described above (28). Twenty-four hours later, reconsolidation was initiated in one group by exposing the rats to an isolated retrieval trial (one tone presentation; Ret group, n = 8), whereas the control group was placed in the same context but was not presented with a cue retrieval (no tone presentation; No Ret group, n = 8). One hour later, extinction training occurred (the No Ret group was presented with 19 CSs and the Ret group with 18 CSs, in the absence of the US; that is, tones were repeatedly presented in the absence of shocks). In the Renewal experiment (Fig. 2), rats were fear-conditioned in context A and then received the retrieval, or context-only exposure, and the extinction session in context B (smooth black floor and peppermint scent) (28). Twenty-four hours later, they were tested for long-term memory in context B, and the next day were tested back in context A (renewal test). We found that the No Ret and Ret rats exhibited similar levels of freezing (a measure of fear expression) during fear conditioning (repeated-measures ANOVA, P > 0.05). No last four trials of extinction (repeated-measures ANOVA, P > 0.1), and at the test of LTM (repeated-measures ANOVA, P > 0.1). When placed back in context A (the original context in which fear to the CS was acquired) to assess whether they would show increased freezing relative to the extinction context (which would be indicative of fear renewal), there was a significant Group × Time of Test interaction, which suggested that the retrieval procedure induced a differential effect on behavior [F(1, 14) = 13.522, P = 0.002]. Follow-up t tests revealed that whereas the No Ret group showed an increase in freezing in context A relative to context B (P = 0.012), the Ret group did not (P > 0.1).

For the Reinstatement experiment, all procedures (described above) were conducted in context A. Twenty-four hours after extinction, rats received five unpaired footshocks and were tested for reinstatement the next day (Fig. 3). The No Ret (n = 8) and Ret (n = 8) groups froze equivalently during conditioning, extinguished at the same rate, and did not differ during the last four trials of extinction (repeated-measures ANOVAs, all tests, P > 0.1). There was a significant Group × Time of Test interaction, which suggested that the retrieval procedure induced a differential effect on freezing behavior [F(1, 14) = 5.456, P = 0.035]. In agreement with previous research, follow-up comparisons revealed that the No Ret rats showed increased freezing 24

---

Fig. 1. Finite lability window to prevent return of fear via post-retrieval extinction. (A) Rats were fear-conditioned (Fear Cond) with three tone-shock pairings. After 24 hours, they were exposed either to an isolated cue retrieval trial (Ret) or context only (No Ret) followed by extinction training. The time interval between the retrieval trial (or context exposure, n = 12) and the extinction was either within (10 min, n = 8; 1 hour, n = 8) or outside (6 hours, n = 8; 24 hours, n = 8) the lability window. Twenty-four hours after extinction, all groups were tested for LTM, and 1 month later for spontaneous recovery. The gray shading represents context A. (B) All groups were equivalent for the last four trials of extinction and at the 24-hour LTM test. One month later, the Ret groups with an interval outside the reconsolidation window (gray), as well as the No Ret group (black), showed increased freezing (spontaneous recovery) relative to the 24-hour LTM test [no retrieval, P < 0.0001; 6-hour interval interval (11), P = 0.001; 24-hour interval (11), P = 0.031]; however, the groups with an interval within the lability window (red) did not (10 min, P = 0.888; 1 hour, P = 0.156) (28). All data points show means ± SEM. Asterisk denotes significance at the 0.05 level.

---

Fig. 2. Attenuation of fear memory by presenting a single isolated retrieval trial followed by an extinction session prevents renewal. (A) Rats were fear-conditioned in context A. Twenty-four hours later, they were exposed either to an isolated cue retrieval trial (Ret, n = 8) or context only (No Ret, n = 8) in context B, followed 1 hour later by extinction training in context B. Twenty-four hours after extinction, they were tested for LTM in context B. The gray shading represents context A; the blue shading represents context B (28). (B) Rats from both experimental groups froze equivalently during the LTM test (all ANOVAs, P > 0.1). When they were placed back in the acquisition context, the No Ret group (black) showed fear renewal (P = 0.012), but the Ret group (red) did not (P > 0.1), relative to their respective LTM tests. All data points show means ± SEM. Asterisk denotes significance at the 0.05 level.
hours after the unsignaled footshocks (reinstate-
ment) relative to the last four trials of extinction
(P = 0.017), but rats in the Ret group did not (P >
0.1). There was no difference between the groups
in pre-CS freezing (fig. S1).

We next proceeded to determine what molec-
ular mechanism might account for the clear
behavioral effect of presenting a single isolated
retrieval trial before extinction training. We
wanted to use a design that would allow us to
examine acute retrieval-induced biochemical
changes that would be taking place on a brief
time scale, but that would also be predictive of
long-term synaptic plasticity, because there is an
overlapping locus of plasticity for extinction and
fear conditioning in the lateral amygdala (29). At
initial retrieval (first CS presentation after condi-
tioning), both reconsolidation and extinction
mechanisms are engaged (3). Generally, as more CSs
are presented, learning becomes biased toward
extinction. In the current study, the only differ-
ence between our two experimental groups for
the behavioral experiments was the interval be-
tween the first and second CSs. For these reasons,
our hypothesis was that a different mechanism
must be engaged early on (at the time of our
differential manipulation), and that this would
lead to a different long-term outcome. It was pre-
viously shown that increasing cAMP-dependent
protein kinase (PKA) signaling facilitates, and its
blockade hinders, reconsolidation of fear memo-
ries in rats (7). In addition, recent rat studies show
that reconsolidation after retrieval of a fear mem-
ory requires phosphorylation of Glur1 glutamate
receptors at the PKA site (Ser445) (30). Phar-
mylation at the Ser445 site is usually followed by
Glur1 receptor insertion (30). Glur1 receptor
insertion is indicative of synaptic plasticity and
takes place during consolidation of fear memo-
ries (31). In addition, Hu and colleagues (32)
recently showed that norepinephrine [which is
known to be important for reconsolidation of
fear memories in both rats (33) and humans (1)]
can trigger Glur1 phosphorylation via PKA
(32). In the final experiment, we therefore ex-
amined the effect of an isolated retrieval on the
phosphorylation of Glur1 at Ser445 and then
tested what the effect of a subsequent CS pre-
sentation would be.

We examined the effect of a single CS
presentation on Glur1 phosphorylation 3 min
and 1 h after the retrieval cue, and then
explored what would happen if another CS
was played 3 min versus 1 h after (Fig. 4).
These time points were chosen because our
two experimental groups (No Ret and Ret) show
a drastically different behavioral outcome, and
their only distinguishing characteristic is a dif-
ferent interval between the first and second CS.
We hypothesized that a certain time period
might be necessary for the memory trace to be
destabilized. Rats were fear-conditioned, then 24
hours later received (i) context exposure only (No
CS) and euthanized 3 min later (n = 6); (ii) a single
CS retrieval and euthanized 3 min later (n = 4); (iii)
a single CS and euthanized 1 hour later (n = 6); (iv)
two CSs with a 3-min interval and euthanized 3 min
later (n = 6); or (v) two CSs with a 1-hour interval and
euthanized 3 min later (n = 6). (B) Quantification
showing an increase in Glur1 phosphorylation at Ser445
[ omnibus ANOVA across all groups, P < 0.05,
with significant post hoc comparisons (Tukey) be-
 tween the CS–3 min and No CS groups, P <
0.05, and between the CS–1 hour and No CS
groups, P < 0.05]. A second CS presented
1 h after initial retrieval resulted in dephos-
phorylation of Glur1 within 3 min, possibly sug-
gesting destabilization of the memory trace, and
may underlie the lack of fear reemergence ob-
served in our behavioral experiments. This
dephosphorylation of Glur1 was not simply due
to the presentation of two CSs instead of one,
because the presentation of two CSs with the 3-
min interval used in standard extinction did not
result in dephosphorylation of Glur1 (Fig. 4 and
fig. S2). These results were also confirmed by
enzyme-linked immunosorbent assay (ELISA).

Fig. 3. Presenting a single isolated retrieval trial before an extinction session prevents reinstatement. (A)
Rats were fear-conditioned. The next day, they were exposed either to an isolated cue retrieval trial (Ret, n = 8) or context only (No Ret, n = 8), followed 1 hour later by extinction training. Twenty-four hours after
extinction, they received five unsignaled footshocks, and the next day were tested for reinstatement.
The gray shading represents context A. (B) The No Ret and Ret groups froze equivalently to the last four CSs of
extinction; however, 24 hours after the unsignaled footshocks, the No Ret group (black) showed increased
freezing (reinstatement) (P < 0.05), but the Ret group (red) did not (P > 0.05). All data points show means ±
SEM. Asterisk denotes significance at the 0.05 level.

Fig. 4. Dephosphorylation of Glur1 Ser445 underlies destabilization and allows behavioral updating during
reconsolidation. (A) Rats were fear-conditioned, then 24
hours later received (i) context exposure only (No
CS) and euthanized 3 min later (n = 6); (ii) a single
CS retrieval and euthanized 3 min later (n = 4); (iii)
a single CS and euthanized 1 hour later (n = 6); (iv)
two CSs with a 3-min interval and euthanized 3 min
later (n = 6); or (v) two CSs with a 1-hour interval and
euthanized 3 min later (n = 6). (B) Quantification showing
an increase in Glur1 phosphorylation at Ser445
[ omnibus ANOVA across all groups, P < 0.05,
with significant post hoc comparisons (Tukey) be-
 tween the CS–3 min and No CS groups, P <
0.05, and between the CS–1 hour and No CS
groups, P < 0.05]. A second CS presented
1 h after initial retrieval resulted in dephos-
phorylation of Glur1 within 3 min, possibly sug-
gesting destabilization of the memory trace, and
may underlie the lack of fear reemergence ob-
served in our behavioral experiments. This
dephosphorylation of Glur1 was not simply due
to the presentation of two CSs instead of one,
because the presentation of two CSs with the 3-
min interval used in standard extinction did not
result in dephosphorylation of Glur1 (Fig. 4 and
fig. S2). These results were also confirmed by
enzyme-linked immunosorbent assay (ELISA).
Fig. 5. Presenting a single isolated retrieval trial before an extinction session leads to less fear memory savings than extinction alone. (A) On day 1, rats were fear-conditioned. The next day, they received either No Ret + Ext (n = 10) or Ret + Ext (n = 10), with a 1-hour interval between the retrieval and extinction phases. Then, on the third experimental day, rats were reconditioned using a single CS-US pairing. The fourth day, we tested the groups and compared them for savings. (B) The No Ret group froze significantly more than did the Ret group during the LTM-savings test presented 24 hours after the single CS-US pairing session (F(1,18) = 11.679, P = 0.003). The No Ret and Ret groups did not differ during extinction (P > 0.1), nor during the single CS-US pairing session (P > 0.1), and no significant pre-CS freezing was observed. All data points show means ± SEM. Asterisk denotes significance at the 0.05 level.

Fig. 6. An isolated retrieval trial followed by an extinction leads to a revaluation of the stimulus as safe, and retards subsequent acquisition of fear conditioning. (A) On day 1, rats were fear-conditioned. On day 2, they received either a retrieval (Ret + Ext, n = 9) or not (No Ret, n = 14), followed 1 hour later by an extinction session (18 CSs for the Ret group, 19 CSs for the No Ret group). On day 3, we reconditioned these groups, as well as conditioned a naïve group of rats (control, n = 7), using five CS-US pairings, to look at the effect of our treatment on reacquisition. (B) The isolated retrieval presented before extinction (Ret + Ext, red) retards reacquisition, relative to a naïve group (white) or the No Ret + Ext group (black). Repeated-measures ANOVA revealed a main effect of Group (F(1,27) = 85.85, P < 0.0001); and a Group × Trial interaction (F(2,27) = 55.687, P = 0.016). Simple main effect follow-up showed that the Ret + Ext group was significantly lower than the Control (P = 0.019) and No Ret (P = 0.009) groups. The Control and No Ret groups were not significantly different from one another. All data points show means ± SEM. Asterisk denotes significance at the 0.05 level.

These findings suggest that the two different treatments (Ret + Ext versus No Ret + Ext) engage different molecular mechanisms in the lateral amygdala and lead to a drastically different behavioral outcome.

To better address whether our Ret + Ext paradigm led to a permanent revaluation of the CS, we next sought to examine subsequent susceptibility to reconditioning. We performed a savings experiment, in which the initial phases were identical to the ones presented in the initial set of experiments (day 1: conditioning with three CS-US pairings; day 2: Ret + Ext, or Ret + Ext, with a 1-hour interval between the retrieval and extinction phases). Then, on the third experimental day, we reconditioned rats through a single CS-US pairing. One additional group received the single CS-US pairing only. On the fourth day, we tested the groups and compared them for savings (six CS presentations). We found that the No Ret group froze significantly more than the Ret group during the LTM test presented 24 hours after the single CS-US training session (F(1.18) = 11.679, P = 0.003) (Fig. 5). The No Ret and Ret groups did not differ during extinction or during the single CS-US pairing session (P > 0.1). These results could suggest that the initial memory has been reversed (deconsolidated), and/or that the valence associated to the CS has been permanently revalued and reencoded as safe.

To address this further, we ran one additional experiment examining the effect of our Ret + Ext manipulation on the rate of fear reacquisition. On day 1, rats were fear-conditioned using three CS-US pairings. On day 2, they received a retrieval (Ret + Ext, n = 9) or not (No Ret + Ext, n = 14), followed 1 hour later by an extinction session (18 CSs for the Ret group, 19 CSs for the No Ret group). On day 3, we reconditioned these groups and also conditioned a naïve group of rats (control, n = 7), using five CS-US pairings, to look at the effect of our treatment on reacquisition. Our results suggest that the Ret + Ext treatment not only does not lead to savings, it actually retards reacquisition, relative to a group being conditioned for the first time (control) or the No Ret + Ext group undergoing conditioning (Fig. 6).

Taken together, our renewal, spontaneous recovery, reinstatement, and savings experiments point to a rather resilient decrease in fear induced by our Ret + Ext paradigm. Our GluR1 results suggest that a process taking place in the lateral amygdala may underlie this effect. Furthermore, the reacquisition experiment suggests not only that the CS no longer induces a fear response, but that it may now act as an inhibitor (similar to what we might expect from a latent inhibition paradigm). This could mean that interference during reconsolidation led to a progressive deconsolidation of the memory followed by the learning of a new interpretation of the CS, or that during reconsolidation, the new valence associated with the CS was incorporated in the updating. In either case, the initial valence conferred
by the first conditioning session no longer seems to exist in its original fear-inducing form.

In considering clinical implications, it will be important to pursue further what might underlie the retardation of reacquisition induced by our behavioral procedure because it could, in principle, result in maladaptive behaviors in some cases. Future experiments will determine whether a once fear-inducing stimulus can become simply neutral, without necessarily turning it into a safety signal. That is, it remains to be established whether the process described here involves destabilization, deconsolidation, and updating as safe, or simply destabilization and updating as safe during reconsolidation. Future studies should disambiguate “fear expression” from “fear memory” in response to our procedure, and determine the effects on other fear-related assays. Extending these findings to humans would be particularly useful in addressing questions pertaining to the subjective experience resulting from the updated stimulus.

We have shown that presenting extinction training within a reconsolidation window opened by an isolated CS prevents renewal, reinstatement, and spontaneous recovery of fear memory. This suggests that a post-consolidation behavioral manipulation can render a memory labile and rewrite and/or update it. In rodents, manipulating the intertrial interval of CS presentations during extinction (e.g., using massed versus spaced training) has been reported to yield differential effects on extinction (34), but although massed training is better in the short term, it worsens the long-term outcome (35). Thus, an important aspect of the current procedure in preventing the return of fear is that the initial CS be isolated from subsequent ones. It was also recently shown that extinction training applied shortly after fear conditioning can prevent memory consolidation and the return of fear (36). However, subsequent experiments, in both rats and humans, that used variations of this protocol have met with limited success (37, 38). This indicates that the contingencies that function to prevent fear reemergence, either in the context of consolidation or reconsolidation, may be sensitive to subtle manipulations. Our results are consistent with the idea that an adaptive purpose of reconsolidation is to incorporate new information at the time of retrieval, and to update a memory (4, 7, 13)—in the present case leading to destabilization of the initial trace in the lateral amygdala, and the reencoding of the once fear-inducing CS as safe.

References and Notes
28. See supporting material on Science Online.
39. Supported by NIH grants R01 MH083774, P50 MH58911, RO1 MH046156, and ROI MH50104 (J.E.L.); NIH grants N0530627 and N054738 (E.K.); postdoctoral fellowships from the Natural Sciences and Engineering Research Council of Canada, Canadian Institutes of Health Research, and Alberta Heritage Foundation for Medical Research (W.-H.M.); and NIH doctoral fellowship F33MH083472 (K.C.C.). We thank K. Nader, M. Davis, C. Cain, J. Johansen, D. Schiller, and G. Quirk and members of his lab for helpful comments.

Supporting Online Material
www.sciencemag.org/cgi/content/full/316/7975/DC1
Materials and Methods
Figs. 1 to 53
3 November 2008; accepted 23 March 2009
Published online 2 April 2009; 10.1126/science.1167975
Include this information when citing this paper.