A growing body of evidence suggests that sleep plays a role in memory consolidation, although the exact nature of this relationship remains unclear. In human studies, sleep-deprived subjects have impaired memory retention (Bonnet and Arand 1995; Arnold et al. 1997; Carter et al. 2003). This impairment, however, may be caused by non-sleep factors associated with the method of deprivation, such as stress. Sleep deprivation in humans can cause physiological stress, marked by stomach ulcers and elevated cortisol (Mullington et al. 2009), that could interfere with memory (Sapolsky 2004). Moreover, the fatigue associated with sleep deprivation may produce performance deficits, even after a night of recovery sleep. One recent imaging study found that it takes more than one night of recovery sleep for the human brain to return to normal use of its neural network (McKenna et al. 2009). This suggests that sleep deprivation produces changes in brain state that could impair normal consolidation or retrieval.

To avoid these confounds, many researchers have turned to a more naturalistic method of allowing normal sleeping or waking periods to pass between training and test. Typically, a sleep group is trained at night and tested in the morning after a period of sleep, while an awake group is trained in the morning and tested in the evening after an equivalent delay interval. Recall is often better after a night of sleep (Ellenbogen et al. 2006; Nishida et al. 2009). This alternative method successfully circumvents the confounds of sleep deprivation; however, most animal studies continue to use sleep deprivation to explore the effects of sleep on memory consolidation.

Early studies deprived animals of sleep by placing them on a rotating disk or on top of an inverted partially submerged flower pot over water (Dement et al. 1968; Van Hulzen and Coenen 1979; Murison et al. 1982; Rechtschaffen et al. 1989). These methods of sleep deprivation have been highly criticized for their potentially stress-inducing side-effects (Horne and McGrath 1984; Smith 1985; Vertes and Eastman 2000). Recent sleep deprivation studies have utilized a gentler approach—handling the animals during sleep. However, even this handling method has been shown to induce stomach ulcers (Murison et al. 1982) and increased levels of glucocorticoids, which can impair cognition (Plihal et al. 1996; Sapolsky 2004). In fact, long-term potentiation is diminished in area CA1 from sleep-deprived rats, and this correlates with increased corticosterone levels (Campbell et al. 2002). Finally, because sleep is a homeostatic drive, the buildup of sleep debt is likely stressful in itself (Roehrs et al. 1990; Carter et al. 2003; Anderson and Horne 2008). Thus, while the rodent sleep deprivation literature suggests that sleep is important for memory consolidation, it is unclear whether these effects are caused by the absence of sleep or by non-sleep-related consequences of deprivation.

An ideal task to examine the relationship between sleep and memory consolidation is Pavlovian fear conditioning, in which a tone is paired with a shock in a distinct context. After a single pairing, rodents will exhibit fear when presented with the training tone, or when returned to the training environment. This latter phenomenon, known as contextual fear conditioning, has garnered considerable interest because it is hippocampus-dependent and has become a prominent rodent model of declarative memory (Anagnostaras et al. 1999, 2000, 2001, 2002a). Pavlovian fear conditioning is well-suited for the examination of the role of sleep and memory because it is rapidly acquired and can dissociate between hippocampus-dependent and independent memory. Consistent with human declarative memory, contextual fear gradually becomes independent of the hippocampus, as this memory is consolidated to neocortical structures (Maren et al. 1998; Anagnostaras et al. 1999; Frankland et al. 2004; Quinn et al. 2008). This consolidation process is thought to reflect coordinated activity whereby fast-changing connections in the hippocampus initially subserve the memory, and over time entrain slow-changing connections in the neocortex, at which time the hippocampus is no longer necessary to maintain the memory (Squire and Alvarez 1995). In contrast, tone (cued) fear is independent of the hippocampus (Anagnostaras et al. 2001), yet both contextual and cued fear memory depend on the amygdala for the animal’s lifetime (Gale et al. 2004).

Graves and colleagues examined the effects of sleep deprivation on Pavlovian fear conditioning (Graves et al. 2003). They found that sleep deprivation impaired performance only when administered immediately following training. Several issues make their data difficult to interpret. First, all three groups (non-sleep-deprived, sleep-deprived immediately after training, and sleep deprived 5 h after training) were trained and tested in the main sleep period (Fig. 1A). Training the animals during their main sleep period likely induced sleep deprivation in all groups, including the Non-Sleep-Deprived group. Moreover, testing mice during this phase is equally problematic (Chaudhury and Colwell 2002). Second, the animals were tested within 24 h of the sleep deprivation,
and performance deficits could result from residual fatigue (Rickard et al. 2008; McKenna et al. 2009). Third, there was no adequate control for the sleep deprivation method used, to demonstrate that it was lack of sleep per se that produced the deficit. Therefore, it is still unclear whether the impairments observed with sleep deprivation are due to the lack of sleep or to other non-sleep-related effects.

In experiment 1, we expanded on the findings of Graves et al. (2003) and used a design that addresses the confounds listed above, to reexamine how sleep deprivation by gentle handling, and the handling manipulation itself, affects memory consolidation. In experiment 2, we examined an alternative method of sleep deprivation by administering two spaced injections of a moderately high dose of amphetamine.

Subjects
Sixty-seven (Exp 1) and 65 (Exp 2) hybrid C57BL/6j x 129T2SvEms/J (Crawley et al. 1997; Matynia et al. 2008) male and female mice (Jackson Laboratory, Bar Harbor, ME) were balanced across groups. Mice were entrained to a 12:12 light/dark cycle. All animals were handled five times for 1 min each in both the dark and light phases.

Experiment 1
Pavlovian fear conditioning was conducted 1 h before the primary sleep period (Fig. 1B). Mice were placed in a fear conditioning chamber (see Conditioning Contexts in Cai et al. 2009; Wood and Anagnostaras 2009) and, after a 2-min baseline, received one tone (2.8-kHz, 30-sec, 85-dB) footshock (2-sec, 1.0-mA, AC) pairing; they remained there for an additional 5 min extended post-shock freezing test. Mice then received one of three treatments. The Control group was undisturbed in their home cage until testing 72 h later. The Sleep/Handle (Sleep-Deprivation) group was handled for 12 h (the entire awake phase), beginning immediately post-training. The Awake/Handle group was handled for 12 h (the entire awake phase), starting 12 h following training. (C) Schematic timeline of experiment 2. Mice were trained at 6 a.m., 1 h prior to their main sleep phase, and tested 72 h later. The Sleep/Sal group was administered saline immediately post-training and 4 h later. The Sleep/Amph group was sleep-deprived with 8 mg/kg (i.p.) of amphetamine immediately post-training and 4 h later. To control for non-sleep-related effects of amphetamine, the Awake/Sal and Awake/Amph groups were administered saline and amphetamine, respectively, 12 h and 16 h post-training.
(Awake/Handle) showed no deficit (Fig. 2C). Freezing during the 2-min baseline was subtracted from the average freezing to the three tones. A one-way ANOVA revealed group differences \((F(2,64) = 3.97, P = 0.02)\), and post-hoc analysis using Fisher’s PLSD revealed a significant difference \((P < 0.05)\) between the Control group and the Sleep/Handle group. No difference was found between the Control group and the Awake/Handle group \((P > 0.8, n.s.)\). Overall, handling during the sleep phase produced pervasive deficits in contextual and tone fear. On the other hand, handling during the awake phase produced selective deficits in contextual fear.

Experiment 2

Training was identical to Exp 1, although a 4-min baseline period was used. After training, mice were given two injections of 8 mg/kg D-amphetamine hemisulfate (Sigma-Aldrich), i.p., or 10 mL/kg saline solution (Fig. 1C). The Sleep/Amph (sleep-deprived) group received amphetamine immediately after training, at the start of their main sleep phase, and 4 h later. The Sleep/Sal control group received saline at the same times. To control for effects of amphetamine, the Awake/Amph and Awake/Sal groups were administered amphetamine and saline, respectively, at the start of the main wake phase and 4 h later. A two-way ANOVA revealed no group differences during training \((F(2,64) < 1, n.s.)\), and all groups exhibited significant learning \((F(8,64) = 17.5, P < 0.001; \text{Fig. 3A})\).

Contextual fear was examined 72 h after training (as in Exp 1). Amphetamine failed to produce any deficit in contextual fear conditioning when given during either the sleep or awake phase (Fig. 3B). A two-way ANOVA revealed no main effect of drug \((F(1,61) < 1, n.s.)\) or phase \((F(1,61) < 1, n.s.)\).

As in Exp 1, a tone test was conducted 30 min after the context test. Amphetamine failed to produce any deficit in cued fear conditioning when given during the sleep or awake phase (Fig. 3C). Freezing during the 2-min baseline was subtracted from the average freezing to the three tones. A two-way ANOVA revealed no main effect of drug \((F(1,61) < 1, n.s.)\) or phase \((F(1,61) = 3.09, n.s.)\). Overall, sleep deprivation by amphetamine had no effect on contextual or cued fear memory.

We found that gentle post-training handling impaired context memory regardless of sleep. These findings indicate that gentle handling produces contextual memory deficits through non-sleep-related factors. Moreover, handling during the sleep phase disrupted hippocampus-independent cued memory. This is in contrast to Graves and colleagues (2003), who found that sleep deprivation administered immediately after training (only) impaired contextual fear, while cued fear was spared. These discrepancies may be due to procedural differences. Graves et al. trained and tested all groups in the middle of the day, which is the main sleep phase for mice (Fig. 1A). Mice were then given no handling ("non-sleep-deprived") or 5 h of handling either immediately after training ("0–5 h") or 5 h later ("5–10 h"). This protocol raises a number of issues. First, all mice experienced some sleep deprivation because sleep was interrupted to give training and testing. Therefore, any sleep deprivation effect observed is additive with a sleep deprivation baseline. Second, deprivation in the 0–5 h group did not span the entire sleep period, nor did it target a specific sleep period, since handling was given 5 h after the sleep phase began. Third, the 5–10 h group did not target sleep or awake periods specifically, as handling overlapped both phases. Fourth, the study did not have a proper control group to examine the effects of handling alone on the memory impairment (i.e., handling only in the awake phase). Due to these confounds, the results of the Graves et al. study are problematic.

Our findings suggest that gentle handling is not a selective method of sleep deprivation. First, extensive handling during the entire sleep phase appears to produce nonspecific impairment. Second, with regard to contextual memory, handling during the awake phase can induce the same magnitude of amnesia as sleep deprivation. This suggests that there are non-sleep-related consequences of extensive handling (e.g., stress, interference, etc.) that may impair memory. This interpretation is consistent with
prior critiques of sleep deprivation studies, which found that seemingly benign handling methods can induce health problems and an elevated stress response (Murison et al. 1982; Mullington et al. 2009). Thus, this common method of sleep deprivation is confounded by non-sleep-related effects and is not an ideal tool for investigating the relationship between sleep and memory.

To explore an alternative to handling, we examined pharmacological sleep deprivation. We administered two high doses (8 mg/kg) of amphetamine immediately after training and 4 h into the sleep phase, and found that this method failed to produce impairments of contextual or cued fear. One possibility for the lack of impairment is that the drug did not actually deprive the animals of sleep. This is unlikely, as this high dose produces insomnia and hyperlocomotion (Ervin et al. 1981; Anagnostaras and Robinson 1996; Anagnostaras et al. 2002b; Wood and Anagnostaras 2009) and D-amphetamine has a long, although variable, half-life (3.5–4.2 h in acidic urine) (Buxton 2006). Another possibility is that amphetamine had a positive effect on consolidation that counteracted the deprivation impairment. This also seems unlikely, as we have found pervasive memory deficits using this dose of amphetamine (alone) during training (Wood and Anagnostaras 2009). Alternatively, these data could indicate that sleep is not involved in memory consolidation. In this manner, stimulant-induced sleep deprivation may be unique in that it does not produce an amnesic stress response. Nonetheless, the unknown effects of amphetamine on stress and consolidation suggest that stimulants are not a viable method to investigate the relationship between sleep and memory.

Sleep deprivation itself may be inherently stressful, perhaps due to a build up of a sleep homeostatic drive. Because sleep is typically treated as a drive (need), governed by homeostatic factors, a “sleep pressure” is thought to build up, which is suggested to be aversive and stressful (Bolles 1967). If this is the case, no sleep deprivation method will satisfactorily dissociate the effects of stress and lack of sleep on cognitive impairment. Therefore, we favor using a more naturalistic paradigm to investigate the consolidation process that may occur during natural sleep. In a separate study published elsewhere, we explored this using a novel rodent paradigm (Cai et al. 2009). We compared 12- and 24-h delay intervals to control for time passage and circadian effects. We controlled for differences in circadian activity by selecting training and testing times within the transition interval from active to wake periods. That study found that sleep plays an important and selective role in contextual fear conditioning, whereby contextual memory is enhanced only when tested after a sleep period. This approach provides a useful way to examine sleep-induced memory enhancement, while avoiding confounds associated with the sleep-deprivation paradigm.

The current data suggest that it is problematic to use sleep deprivation as a way of exploring the role of natural sleep in memory consolidation. Although gentle handling produced amnesia, it was not specific to the sleep phase, making it difficult to disentangle the cause for the memory impairments. Nonetheless, the effects of sleep deprivation may be intrinsically interesting, as sleep deprivation is so prevalent in modern society (Bonnet and Arand 1995; Arnold et al. 1997; Anderson and Horne 2008). However, rather than extensive handling of animals, we would encourage utilization of more naturalistic modeling of sleep deprivation.

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