Cognition and Behavior

Sleep Deprivation Disrupts Acquisition of Contextual Fear Extinction by Affecting Circadian Oscillation of Hippocampal-Infralimbic proBDNF

Wei Sun,1,2 Jia Li,2 Shuai Cui,2 Le Luo,3 Peidong Huang,4 Chunzhi Tang,2 and Lei An1,2,3,5

https://doi.org/10.1523/ENEURO.0165-19.2019

1The First Affiliated Hospital of Guizhou University of Traditional Chinese Medicine, Guiyang 550001, China, 2Medical College of Acupuncture-Moxibustion and Rehabilitation, Guangzhou University of Chinese Medicine, Guangzhou 510006, China, 3College of Acupuncture-Moxibustion and Orthopedics, Guizhou University of Chinese Medicine, Guiyang 550025, China, 4Medical College of Acupuncture-Moxibustion and Rehabilitation, Yunnan University of Traditional Chinese Medicine, Kunming 650500, China, and 5Department of Physiology, University of Saskatchewan, Saskatoon, Saskatchewan S7N 5E5, Canada

Abstract

Extensive evidence showed that mature brain-derived neurotrophic factor (mBDNF) levels displayed a circadian pattern. Circadian disruption, for example, sleep deprivation (SD), induced functional and behavioral deficits. However, compared with that of mature form, the biological role of the pro-peptide, proBDNF, was poorly understood. Here, we found that proBDNF was expressed under circadian rhythm in the ventral hippocampus (vHPC). SD rats exhibited deficits in acquisition of conditioned extinction and damped rhythmicity in vHPC proBDNF activity that were accompanied by SD between zeitgeber time (ZT)0 and ZT4, but not the late stage of sleep period. Furthermore, SD affected fear extinction through vHPC-IL proBDNF signaling, which was associated with NR2B subunits of NMDA receptors. More importantly, infusion of proBDNF could mitigate SD-induced abnormal neural activity, by suppressing the enhanced basal firing rate of IL-RS and elevating the depressed neural response that evoked by acquisition of conditioned extinction. Therefore, this finding provided the first evidence that circadian oscillation of vHPC proBDNF activity contributed to the effects of SD on acquisition of conditioned fear extinction, and suggested a new therapeutic target to reverse the cognitive deficits in sleep-related mental disorder, such as post-traumatic stress disorder (PTSD).

Key words: circadian oscillations; HPC-IL pathway; neural activity; NMDA receptors; sleep deprivation

Significance Statement

The aim of this study was to assess the circadian role of pro-brain-derived neurotrophic factor (proBDNF) in the ventral hippocampus (vHPC) on contextual fear extinction and detect whether sleep deprivation (SD) affected the circadian rhythm of proBDNF levels and attempted to explore the underlying mechanisms of functional and behavioral deficits. These findings provided the first evidence that circadian oscillation of vHPC proBDNF activity contributed to the effects of SD on acquisition of conditioned fear extinction, and offered potential avenues to mitigate sleep-related cognitive and functional disorders, such as PTSD and depression.
Introduction

The neurotrophin brain-derived neurotrophic factor (BDNF) is a member of a family of neurotrophic factors critically involved in various physiologic functions, particularly neuroplasticity, memory and sleep (Nakai et al., 2014; Schmitt et al., 2016), suggesting its biological role throughout life (Lu et al., 2009). BDNF protein is biosynthesized as a precursor of BDNF (proBDNF), which can be secreted and processed extracellularly by plasmin or by matrix metalloproteases to produce mature BDNF (mBDNF; Gray and Ellis, 2008). Recent studies in rodents indicated the endogenous cyclical alterations in central mBDNF, its cognate receptor, and other related candidate effectors during sleep were necessary for learning and memory consolidation (Eckel-Mahan et al., 2008; Jang et al., 2010; Zhang et al., 2013). As the pro-peptide of mBDNF, it is quite possible that proBDNF plays a critical role in the regulation of circadian rhythms.

Circadian rhythms were seen in the sleep and wake cycle as well as in several other aspects of physiologic functioning, such as strengthening and integration of newly formed memories (Eckel-Mahan et al., 2008; Gerstner and Yin, 2010; Michel et al., 2013), including fear memories (Pace-Schott et al., 2015; Boyce et al., 2016). According to the synaptic-homeostasis hypothesis, homeostatic regulation of synapse formation during sleep was attributed to reactivation of the underlying neuronal memory traces (Griffith and Roshbash, 2008; Vyazovskiy et al., 2008). Indeed, wakefulness appeared to be associated with synaptic potentiation, whereas sleep may favor global synaptic depression, thereby preserving an overall balance of synaptic strength (Vyazovskiy et al., 2008). This view was also supported by the prominent neuronal feature of sleep in acquainting new information during the following wake-phase (Marshall et al., 2014; Li et al., 2017).

Accumulating evidence has shown that proBDNF and mBDNF elicited seemingly opposite biological effect on modulating brain structure and function (Holm et al., 2009; Je et al., 2012; Mizui et al., 2015). Recently, there was growing interest on the mechanisms of neurotrophin effects on plasticity underlying fear extinction learning (Cowansage et al., 2010; Andero and Ressler, 2012). Distinct from the enhanced effect of mBDNF on LTP, proBDNF can negatively regulate neural remodeling by facilitating the induction of LTD (Yang et al., 2014), which weakens synaptic transmission to enhance learning (Dalton et al., 2012; Walker et al., 2015) and memory (Dalton et al., 2012; Li et al., 2016) of fear extinction. Considering that the critical role of sleep in extinction memory was associated with BDNF signaling (Hill et al., 2016) and the negative retrograde effect of proBDNF on dendritic complexity and spine density (Yang et al., 2014), synaptic transmission (Yang et al., 2009), and cell survival (Je et al., 2012; Sun et al., 2012) was strongly linked to memory decay, it remained unclear whether sleep-dependent circadian rhythms of proBDNF expression played a pivotal role in fear extinction.

Empirical researches have shown that the amygdala, hippocampus, and mPFC were involved in fear conditioning (Sotres-Bayon et al., 2009; Milad and Quirk, 2012). As one of important BDNF-containing inputs, ventral hippocampus (vHPC) was important for supplying BDNF to the IL mPFC to facilitate context-dependent fear learning (Wiltgen et al., 2006; Peters et al., 2010) and extinction (Corcoran et al., 2005; Sierra-Mercado et al., 2011). At the cellular level, increased excitability in infralimbic cortex (IL-PFC) following extinction has been shown to predict extinction success (Burgos-Robles et al., 2007; Santini et al., 2008). Hippocampal mBDNF infusion increased IL-PFC neuron firing (Rosas-Vidal et al., 2014), suggesting that coordinated mBDNF release in the HPC-mPFC circuit. Therefore, the dysfunction of homeostatic regulation induced by sleep pressure may change the cortical firing correlate of behavioral states. Also unknown is which proBDNF-mediated pathway was mainly correlated with SD-induced fear extinction acquisition?

To address above issues, we detected the circadian variations of proBDNF in the vHPC and the effect of SD on it. By pharmacological infusions, we attempted to verified fear extinction learning was dependent on the intact circadian of vHPC proBDNF expression. Considering the molecular mechanism linking BDNF/TrkB signaling with NMDA receptors in memory (Nakai et al., 2014; Itoh et al., 2016), we also infused antagonist of NMDA-2A or NMDA-2B subunit into vHPC during SD to detect the interaction between proBDNF and NMDA receptors during fear extinction. Finally, by single-unit recording, we inspected that the circadian oscillation of proBDNF level was modified within HPC [or basolateral amygdala (BLA)]-IL circuit. Our findings supported the hypothesis that the circadian oscillation of proBDNF activity in vHPC was critical for fear extinction acquisition, while the interference in the circadian peak of proBDNF expression by SD pointed a core feature of the mental disorder.

Materials and Methods

Animals and drug administration

Three-month-old male Sprague Dawley rats (Beijing Research Center for Experimental Animals) were individually housed in standard Macrolon cages with free access to food and water unless food was restricted to prepare for level press experiment. Animals were kept on a 12/12 h light/dark cycle (L/D; lights on at 8 A.M.) unless otherwise specified. A layer of sawdust served as bedding. During the light phase, stalls were lit by two 200-W fluorescent light bulbs that produced an average light intensity of 300–400 lux at the level of cages. Lux was detected using a light meter (VWR Scientific). Each animal...
was handled extensively (10 min/d) till the experimental day. All experiments were performed according to the ethics Committee on the Care and were approved by the University of Animal Research Ethics Board.

Groups and treatments were indicated in figure and its legend. All control animals were treated with the same dose of saline or artificial CSF (ACSF).

**Sleep deprivation (SD)**

Three to 7 d before being used, all animals were transferred to a testing room controlled for temperature (21–23°C), relative humidity (40–50%), and noise (40–50 dBA during the light phase). Subgroups of rats were regularly monitored for their sleep-wake behavior via the infrared detector (Mini Mitter) was attached to each cage to detect activity. Infrared beam breakage was recorded using VitalView Data Acquisition System (Mini Mitter). Data were transformed into bins per min using ClockLab (Actimetrix). The actograms shown were plotted the locomotion of rats in the corresponding L/D or D/D conditions. The locomotor activity during light or dark phase was measured. All data were normalized by the activity of control group during the light phase.

Rats were subjected to 4-h SD from either zeitgeber time (ZT)0 to ZT4 or ZT4 to ZT8. These time window were chosen dependent on circadian oscillation of proBDNF activity during the light phase (see Results) and the previous findings in sleep’s role in learning and memory (Carter et al., 2009; Yang et al., 2012; Ackermann and Rasch, 2014; Ravassard et al., 2016; Vyazovskiy et al., 2017). For example, early studies indicated that the early period of sleep (such as from ZT0 to ZT4; Carter et al., 2009; Vyazovskiy et al., 2017) was critical for promotion of naturally acquired extinction (Rasch et al., 2007; Yang et al., 2012; Pace-Schott et al., 2015), erasing fear information (Rolls et al., 2013; Ravassard et al., 2016), and enhancing new learning (Rasch and Born, 2013; Stickgold and Walker, 2013; Landmann et al., 2014). However, the second half of the night was important for mirror tracing, priming, and implicit memory (Rasch and Born, 2013; Ackermann and Rasch, 2014). Briefly, animals were kept under constant observation of the experimenter and kept awake by mild sensory stimulation, which involved tapping on the cage, gently shaking the cage or, if necessary, disturbing the sleeping nest, when they assumed a sleeping posture (Hagewoud et al., 2010; Oyanedel et al., 2015). Previous studies have shown that this procedure was effective in keeping rodents awake for several hours, as established by electroencephalic recordings (Oyanedel et al., 2015), without being a major stressor (Tadavarty et al., 2009). The number of stimuli needed to keep the rats awake was recorded.

**Fear conditioning**

Rats were contextual fear conditioned and extinguished in standard operant chambers (Coulbourn Instruments) inside sound-attenuating boxes (Med Associates). The chambers contained two retractable levers located on either side of a central food trough. White cue lights were located above each lever. Delivery of food pellets was accompanied by an auditory stimulus and illumination of a cue light located within the food trough. As previous studies (Paterson et al., 2005; Peters et al., 2010), animals initially received 45-mg food pellets at a 30-s interval for 30-min initial sessions with no requirement to lever press. When rats consumed all pellets during initial stage, they were trained daily 30-min (between ZT3.5 and ZT4.5) sessions with one of two levers extended randomly when the cue light above the level was illuminated. The schedule was progressively changed according to the sequence fixed ratio (FR)-1, FR-15, FR-30, and finally FR-60. The criterion for moving through the sequence was earning 50 pellets within the 30-min session. Rats were tested in a 30-min session till they reached 10 presses per minute on FR-60.
Bilateral microinjection

Surgery was prepared as our reports (An and Sun, 2017, 2018). Briefly, rats were anesthetized with sodium pentobarbital (60 mg/kg, i.p.), placed in a stereotaxic frame (SN-3, Narishige) for surgery under atropine (0.1 mg/kg, i.p.) which help relieve respiratory congestion. Stainless-steel guide cannulae (22 gauge; Plastics One, Inc.) were bilaterally inserted above two of three sites: vHPC (AP: −6.0 mm, ML: ±5.0 mm, DV: 5.4–5.6 mm), BLA (AP: −2.8 mm, ML: ±5.0 mm, DV: 7.4–7.6 mm), and IL-PFC (AP: +2.8 mm, ML: ±3.1 mm, DV: 3.8–4.2 mm; angled at 30°). For IL cannula implantation, the angled approach was used to avoid backflow to PL (Sierra-Mercado et al., 2011). Sterile stainless-steel stylet (30 gauge, 10 mm, Plastics One Inc.) was inserted into guide cannula to avoid obstruction. All rats were allowed to recover for at least 7 d. Infusions were achieved by inserting 30-gauge dummy needles (10 mm, Small Parts Inc.) linked via PE-50 tubing to a microsyringe pump (Harvard Apparatus), extended 1.0 mm beyond the end of the cannulae. Needles were inserted into both cannula then infused cleavage-resistant proBDNF (2 ng/ml), sheep anti-proBDNF antibody (10 μg/μl), mBDNF [1.5 μg/μl; 3-(2-carboxypiperazin-4-yl)propyl-1-phosphonic acid (CPP; 32 ng/μl)], NVP-AAM077 (0.8 ng/μl), Ro25-6981 (2.0 ng/μl), or ACSF into vHPC, BLA, or IL-PFC area was initiated at 0.5 μl/min per side for 2 min. The drugs were purchased from Sigma-Aldrich Chemicals except for cleavage-resistant proBDNF (Alo-mone Labs), anti-proBDNF antibody (R&D Systems), and human mBDNF (R&D Systems). All doses were determined from published studies, which indicated the safety and efficacy of cleavage-resistant proBDNF (Woo et al., 2005; Holm et al., 2009), anti-proBDNF antibody (Fan et al., 2008; Bai et al., 2016), mBDNF (Peters et al., 2010; Rosas-Vidal et al., 2014), CPP (McDonald et al., 2005; Dong et al., 2013), NVP-AAM077 (Fox et al., 2006; Dong et al., 2013), and Ro25-6981 (Fox et al., 2006; Dong et al., 2013). The infusion needles were left in place for a 5-min period to permit diffusion of the drug. One week before commencement of the experiment, habituation sessions were conducted four times for each rat without infusion. Following testing in all conditions, the rats were sacrificed with urethane and the brain was collected in 10% formalin/10% sucrose. The placements of cannulae were used to identify the location of the infusion sites. Only data obtained from rats with correctly inserted needles were included in statistical analysis.

Protein preparations and Western blot analysis

Additional groups of rats were housed in a L/D for at least two weeks before time course experiment. Rats were killed every 4 h during one 24-h period. During the dark cycle, rats were killed under a dim red light (1–2 lux). vHPC, BLA, and IL-PFC were rapidly dissected on ice and snap-frozen in liquid nitrogen and stored at −80°C. Tissues was homogenized in ice-cold lysis buffer (pH 7.4) containing a cocktail of protein phosphatase and proteinase inhibitors (Sigma) to avoid dephosphorylation and degradation of proteins. Protein concentrations were determined using the bicinchoninic acid (BCA) assay. Twenty micrograms (15 μl) of total protein per lane was resolved in 10–15% SDS-PAGE gels followed by electro-transferring to PVDF membranes (Pall). Non-specific binding of antibodies to membranes was blocked with 5% (w/v) non-fat milk for 2 h at room temperature, followed by incubation overnight at 4°C with the primary mouse anti-proBDNF antibody (1:500; Santa Cruz Biotechnology) or rabbit anti-mBDNF antibody (1:5000; Millipore Bioscience Research Reagents). Mouse anti–β-actin (1:20,000; Sigma) was used as an internal control. After rinsing in Tris-buffered saline-Tween 20, the membranes were incubated with horseradish-peroxidase (HRP)-conjugated secondary goat anti-mouse or anti-rabbit IgG (1:1000; Southern Biotechnology Associates) incubated for 2 h at room temperature. After three rinses (10 min in each) in TBS-T buffer, immunoreactivity was detected by ECL Western Blotting Detection kit (CWBio).

Immunohistochemical staining

Rats were anesthetized and transcardially perfused with saline, then fixed with 4% paraformaldehyde in PBS. The intact brains were post-fixed in 4% paraformaldehyde for an additional 2 h, and placed in 30% sucrose cryoprotectant solution for 48 h. Then the brains were embedded in OCT medium and stored at 80°C. Sagittal sections (10 μm thick) were prepared using a Leica microtome (Leica RM2235, Leica Biosystems). In brief, deparaffinized sections were rehydrated through a graded series of ethanol to PBS. Antigen retrieval was performed in citrate buffer (10 mM, pH 6.0) at 95°C for 40 min. Sections were incubated in 3% H2O2 to clear endogenous peroxidase, blocked and incubated overnight with chicken anti-proBDNF antibody (1:1000; Millipore Bioscience Research Reagents) at 4°C. Negative controls were conducted by exchange of primary antibody for PBS. Biotinylated-conjugated anti-chicken IgG (1:1000; Millipore Bioscience Research Reagents) was incubated for 1 h at 37°C and then visualized with DAB (Sigma-Aldrich). Between steps, the sections were washed twice in PBS. Finally, the sections were counterstained with hematoxylin and mounted. Sections were viewed on a Leica microscope and digitized using a charge-coupled device camera (Olympus DP71). Quantification was performed using Image-Pro Plus software. One section of each region was selected from each rat in each group, and five fields per section were taken for analysis. We examined the sum of integrated optical density (IOD) of proBDNF-positive neurons in each field of a selected section.

Single-unit recording

Microelectrode array was arranged in a 4 × 8 matrix using 25-μm-diameter tungsten wires (California Fine Wires) in a 35-gauge silica tube (World Precision Instruments). It was then attached via gold pins to an EIB-36-P TB board (Neuralynx Inc.). The electrode tips were gold-plated to maintain the impedance to 200–600 kΩ measured at 1 kHz. Rats were anesthetized with isoflurane and prepared for surgery using previously reported procedures (An and Sun, 2018; Sun et al., 2018a). Cannula were implanted...
bilateral into vHPC (AP: −6.0 mm, ML: ±5.0 mm, DV: 5.4–5.6 mm). Electrode arrays were unilaterally implanted into IL-PFC (AP: +3.0 mm, ML: ±0.6 mm, DV: 4.5–5.0). The hemisphere was implanted randomly but counterbalanced across rats of groups. A stainless-steel wire was used as ground electrode and soldered onto one jewelers’ screw, which was implanted into the skull. The electrode array was fastened to the cranium by dental acrylic with additional scull screws as anchors.

During the whole test, nine rats were assigned to one of three conditions (e.g., SD, CON, SD-proBDNF) in each recording session. This was assigned in a pseudorandom order between subjects and treatments during each training day. The recording was conducted for 5 min in the fear conditioning chamber immediately following the extinction training with a Digital Cheetah system (Cheetah software, Neuralyx Inc.). Additionally, a 30-min baseline recording was performed in the fear conditioning chamber 30 min before fear extinction training. Unit signals were recorded via a HS-36 unit gain headstage (Neuralynx Inc.) mounted on animal’s head by means of lightweight cabeling. Unit activity was amplified (1000–10,000 times), sampled at 32 kHz and 600- to 6000-Hz bandpass filters. To verify the stability of recording, unit activities were recorded for ~15 min before the training initiation.

After the completion of all recording sessions, electrolytic lesions (10-µA current for 10 s) were made to mark the selective recording sites, which were identified using standard protocols with reference to The Rat Brain in Stereotaxic Coordinates (Paxinos and Watson, 1997). Only data from rats with probes contained within IL-PFC were analyzed.

Unit isolation and classification criteria

Spike sorting was performed with offline Neuralynx’s software (SpikeSort 3D), using a combination of KlustaKwik, followed by manual adjustment of the clusters (Klusters software package). Briefly, multiple parameters were used to effectively visualize clusters with the most often used combination of spike height, trough, and energy, associated with the waveforms (Insel and Barnes, 2015; Kim et al., 2016). Each cluster was then checked manually to ensure that the cluster boundaries were well separated and wave form shapes were consistent with action potentials (Fig. 6C). Interspike interval histograms were additionally examined for ensuring single unit activity (Fig. 6B). Using methods described elsewhere (Insel and Barnes, 2015; Kim et al., 2016; An et al., 2018; Sun et al., 2018b), units were then graded for quality and classified as regular-spiking projection neuron (RS) and fast-spiking inhibitory neuron (FS) from PFC recording, respectively.

Code accessibility

All code is available as extended data. Code for data processing and analysis can be found at http://neurosuite.sourceforge.net/. We used SpikeSort 3D, Neuroexplorer and MATLAB software for data processing. These were run in Windows 7 OS. All Data are available from the corresponding author upon reasonable request.

Statistical analysis

Data are expressed as mean ± standard error of the mean. The analyses were performed with Neuroexplorer (Nex Technologies), MATLAB (MathWorks), and SPSS 17.0 software. The data of fear conditioning test were compared using a repeated-measures ANOVA as a two-way design in which testing block (block 1, block 2, block 3, block 4, block 5, or block 6) was a within subjects measure, and treatment (control, SD, or other treatments) was a between subjects measure. A one-way ANOVA examined the effects of treatment (control or SD) on the data of open field and motivation tests. The data of Western blot and immunohistochemical tests were subjected to a two-way ANOVA in which brain region (IL-PFC, BLA, or other brain areas) and treatment (control, SD, or other treatments) were dependent variables. To measure the correlation between average firing rate and the ratio of the spike wave form, Pearson’s correlation analysis was performed on unit classification. For comparison of the mean firing rate in the electrophysiological recording, a repeated-measures ANOVA was employed to examine the effects of testing block (block 1, block 2, block 3, block 4, or block 5) as a within subjects measure, and the effects of treatment (control, SD, or SD plus proBDNF treatment) as a between subjects measure. When the ANOVA revealed a significant main effect or interaction between main factors, data were further analyzed by Tukey’s post hoc tests; p < 0.05 level, p < 0.01 level, and p < 0.001 level of confidence were used in the analyses.

Results

SD inhibited proBDNF activity rise during circadian cycle

To determine whether rats entrained to a L/D have oscillations in hippocampal proBDNF activity, vHPC tissues were taken every 4 h starting at light onset (ZT0). During the dark cycle, rats were killed under red light to avoid the influence of light-induced activation on neurotrophin signaling (Chang et al., 2014). As shown in Figure 1, actograms generated from monitoring voluntary activity of rats in L/D condition (Fig. 1A, left) indicated that locomotion rhythms of L/D rats maintained a circadian oscillation (Fig. 1E). Western blot analysis showed a pronounced oscillation in proBDNF expression over the 24-h period (Fig. 1A, right). Since the expression of proBDNF was significantly higher at ZT4 compared with ZT16 (p < 0.001), ZT4 and ZT16 were used as time points for subsequent comparison (Fig. 1F). To verify that the L/D oscillations in proBDNF activity were truly circadian, we tested whether or not they persisted under free-running (D/D) condition (Fig. 1B). Mice were entrained to an L/D cycle for two weeks and then placed in complete darkness (D/D) for at least two weeks before being analyzed. Similar actograms were observed (Fig. 1B, left), locomotor activity of L/D rats was comparable to that of D/D rats (Fig. 1E). As expected, in the D/D condition, vHPC proBDNF at circadian times (CT) 0, 4, 8, 12, 16 and 20 maintained circadian variations (Fig. 1B, right), which were in a similar pattern as the L/D rats’ (Fig. 1F). No sign of circadian rhythmicity in proBDNF activity exhibited in
**Figure 1.** SD interrupted circadian oscillations in proBDNF activity in the hippocampus. Rats were entrained to a standard L/D condition (A), 12/12 h dark/dark cycle (D/D condition; B), partial SD between ZT0 and ZT4 [SD(0–4); C], or partial SD between ZT4 and ZT8 [SD(4–8); D]. Actograms of voluntary movement during L/D (free-running) condition (A, left), D/D (free-running) condition (B, left), SD(0–4) condition and SD(4–8) condition (D, left) showed a circadian oscillation. Each actogram was from individual animal; n = 6–8 per group. The expression of vHPC proBDNF, normalized to actin, during the 24-h cycle in L/D (free-running) condition (A, right), D/D (free-running) condition (B, right), SD(0–4) condition (C, right), and SD(4–8) condition (D, right); n = 7–8 per time point. **E,** When ZT were blocked into 12-h increments, the movement, normalized by the activity of L/D condition during ZT0 and ZT12, showed no different among all conditions; n = 6–8 per group. **F,** Quantification of proBDNF relative to actin protein expression at ZT4 and ZT16; *p < 0.01; n = 7–8 per time point.
striatal or cerebellar tissue (data not shown), indicating either that there was no proBDNF oscillation in striatal or cerebellar neurons, or that there was no synchronized activity in these regions.

SD was conducted on rats between ZT0 and ZT4 [SD(0–4)] to seek whether SD could perturb the circadian variations in proBDNF activity (Fig. 1C). Although the locomotion activity of SD(0–4) rats during the SD period was slightly higher than rats in the L/D and D/D condition, no statistical difference was found (Fig. 1E). Strikingly, proBDNF activity of SD(0–4) rats lost marked 24-h variations in the L/D condition of the day following SD (Fig. 1C, right) and the expression at ZT4 was significantly lower than other groups (all comparisons; p < 0.01; Fig. 1F). To determine the specific time effect of SD on circadian oscillations in proBDNF activity, we performed SD between ZT4 and ZT8 [SD(4–8)], which was the decaying period in circadian variations of proBDNF activity. However, SD from ZT4 to ZT8 did not affect rats’ voluntary movement during L/D condition (Fig. 1D, left) and the synchronized oscillations in proBDNF expression (Fig. 1D, right, F) on the following day. Additionally, there was no difference between SD(0–4) and SD(4–8) conditions in the number of stimuli to keep the rats awake (data not shown), suggesting that the stimulant effects of tapping or shaking were not attributed to the circadian variation.

**SD affected extinction of conditioned fear, but not acquisition, of fear conditioning**

We also tested whether the effect of SD on proBDNF oscillations affected fear conditioning, including extinction [post-COND-SD(0–4) (Fig. 2A, top); post-COND-SD(4–8) (Fig. 2A, middle)] and acquisition [pre-COND-SD(0–4); Fig. 2A, bottom]. Rats trained for contextual fear were sleep deprived between ZT0 and ZT4 on the following day, and then tested extinction of conditioned fear 20 h (pre-extinction at ZT24), 48 h (light phase at ZT4), 60 h (dark phase at ZT16), or 72 h (light phase at ZT4) post-training. No group difference was observed when fear extinction training was conducted before SD [effect of treatment: F(1,8) = 1.56; p > 0.05; the first subset of post-COND-SD(0–4) in Fig. 2C]. Similarly, there was no statistical difference in the freezing levels between CON and SD groups in the Post-COND SD(4–8) or Pre-COND condition (data not shown). However, SD(0–4) rats tested during both light [ZT4; effect of treatment: F(1,14) = 29.72, p < 0.001; block 3–5: p < 0.001; the second subset of post-COND-SD(0–4) in Fig. 2C] and dark [ZT16] phases [effect of treatment: F(1,12) = 43.10, p < 0.001; block 3–5: p < 0.001; the third subset of post-COND-SD(0–4) in Fig. 2C] had deficits in extinction training compared with their matched control rats. This effect can be persistently tested on the third day following conditioning training [F4(30) = 38.86, p < 0.001; block 3–5: post hoc p < 0.001; the last subset of post-COND-SD(0–4) in Fig. 2C], predicting the persistent effect of SD on extinction acquisition. Furthermore, SD during ZT4 and ZT8 following conditioning training did not damage acquisition of fear extinction, which tested 48 h [effect of treatment: F(1,12) = 1.12, p > 0.05; the subset of post-COND-SD(4–8) in Fig. 2C], 60 h [effect of treatment: F(1,12) = 1.33, p > 0.05] or 72 h [effect of treatment: F(1,12) = 1.17, p > 0.05] after conditioning training. When SD was conducted 24 h before the acquisition of fear conditioning, no difference was found during acquisition [effect of treatment: F(1,22) = 0.83, p > 0.05; Fig. 2B] or extinction training that tested at 48 h [ZT4; effect of treatment: F(1,10) = 0.94, p > 0.05; the last subsets of pre-COND-SD in Fig. 2C], 60 h (ZT16; effect of treatment: F(1,10) = 1.29, p > 0.05) or 72 h (ZT4; effect of treatment: F(1,10) = 1.31, p > 0.05). Additionally, the effects of SD on locomotion, anxiety, and motivation behavior were tested 24 h after SD (Fig. 2D). No statistical difference was found in the total travel distance (Fig. 2E) and the percentage time in the center of the apparatus (Fig. 2F) in the open field test, or the motivation behavior (Fig. 2G). Therefore, the alternation in freezing behavior was not attributed to the effect of SD on locomotion, anxiety or motivation defects.

Furthermore, rats were SD between ZT0 and ZT4 immediately following conditioning training (Fig. 3A). We found SD severely disturbed memory consolidation of conditioned fear [effect of treatment: F(2,18) = 24.58, p < 0.001; post hoc p < 0.01; Fig. 3B]. This deficit cannot be reversed by the infusion of proBDNF (post hoc p > 0.01; Fig. 3B). The immunohistochemical staining with anti-proBDNF antibody was conducted 24 h after extinction training. Compared with SD group, the expression of proBDNF was significantly elevated in the brain regions of control group (Fig. 3C), including vHPC, PFC, striatum, cerebellum, and diencephalon (for all comparisons p < 0.05; Fig. 3D), which indicated that proBDNF acts at different brain areas during the consolidation process. However, no statistical difference was found in the amygdala. Meanwhile, a positive correlation between the level of proBDNF in the vHPC and fear behavior was found (Pearson’s r = 0.216; p < 0.001). Together with the finding of circadian variations of proBDNF activity, our results implied the correlation between the oscillations of proBDNF activity in the vHPC and fear extinction, but not initial formation or consolidation fear memory.

**SD impaired circadian proBDNF level through NR2B-containing NMDA receptors**

To verify the effect of the circadian proBDNF level on fear extinction and whether exogenous proBDNF could mitigate extinction deficits induced by SD, rats were subjected to contextual fear conditioning and, the following day, received bilateral vHPC infusion of cleavage-resistant proBDNF, anti-proBDNF antibody or mBDNF before the extinction training day (Fig. 4A). As before, SD rats (SD) showed significantly higher freezing levels relative to control (CON) rats [effect of treatment: F(4,39) = 113.53, p < 0.001; block 3–5: p < 0.001; Fig. 4B]. Additionally, since the experimental procedures were different for the data shown in Figure 2 (three controls groups) and Figure 4, the freezing levels, which were tested 48 h after conditioning training, were qualified. However, no statistical difference was found between the SD groups of Figures 2, 4 (repeated-measures ANOVA, effect of treatment: F(3,23) = 0.96, p > 0.05) or between CON groups of
Figures 2, 4 (repeated-measures ANOVA, effect of treatment: $F_{(3,23)} = 0.73, p > 0.05$). VHPC infusion of proBDNF into SD rats was able to enhance fear extinction learning compared to SD (block 4: *post hoc* $p < 0.01$; block 5 and 6: $p < 0.001$; Fig. 4B). However, infusion of proBDNF antibody into control (CON-anti) rats was indistinguishable from CON rats (block 4–6: $p < 0.001$; Fig. 4B) and SD-proBDNF in their freezing levels (block 4–6: $p < 0.001$; Fig. 4B). Since previous studies found infusion of mBDNF into the IL mPFC promoted extinction memory of conditioned fear even in the absence of extinction training (Peters et al., 2010) and rescued the late-phase of long-term potentiation as well as the amnesia caused by protein synthesis inhibitors (Bekinschtein et al., 2008), firstly, in a parallel study, we have demonstrated that increased mBDNF exhibited the circadian oscillation in the vHPC (data not shown), which was consistent with the reports from other labs (Eckel-Mahan et al., 2008); and then
intra-hippocampus injection of mBDNF was performed during SD (SD-mBDNF) to detect the effect of mBDNF on extinction acquisition. Although SD-mBDNF group learned extinction faster than SD group (block 5 and 6: \( p < 0.01 \); Fig. 4B), it was still slower than CON and SD-proBDNF groups (block 5 and 6: \( p < 0.01 \); Fig. 4B). However, infusion of cleavage-resistant proBDNF into the vHPC at ZT0 could not ameliorate the consolidation defects of SD rats (effect of treatment: \( F_{(2,18)} = 24.58, p < 0.001 \); post hoc \( p < 0.01 \); Fig. 4B). It suggested the impairment of consolidation process was not attributed to the disruptive effect of SD on circadian proBDNF expression. Together all, these results demonstrated that acquisition of extinction was associated with circadian proBDNF oscillation in the vHPC.

Growing evidences suggested extinction memory was linked with the activation of NMDA receptors (Burgos-Robles et al., 2007; Sotres-Bayon et al., 2009). For example, extinction of inhibitory avoidance learning was blocked with a variety of kinase, NMDA, mRNA, and protein synthesis inhibitors into the dorsal HPC (Vianna et al., 2001; Cammarota et al., 2003). Other studies have demonstrated that proBDNF directly or indirectly modified NMDA receptors. Early evidence implied the role of postsynaptically BDNF increased NMDA receptor activity through phosphorylation and upregulation of NMDA receptor subunits (Lin et al., 1998). Moreover, the proBDNF-p75NTR signaling was important for the expression of NR2B-dependent LTD (Woo et al., 2005), which was required for the extinction of conditioned fear and taste aversion (Dalton et al., 2012; Li et al., 2016). More importantly, it has been suggested that the impairment in synaptic plasticity following SD may be due to a reduction in hippocampal levels of BDNF (Schmitt et al., 2016), alternation in NR2A/NR2B ratio of HPC NMDARs (Kopp et al., 2006) and without alternation receptor subunit composition (McDermott et al., 2006). Strikingly, this change was reversed after recovery sleep (Kopp et al., 2006). It was possible, therefore, that the disruption of circadian proBDNF oscillation in SD group, which showed extinction deficits, involved in NMDA receptors signaling. To test this hypothesis, fear-conditioned rats were sleep deprived 30 min preceded by intra-hippocampal infusion combinations of saline and saline (Sal-Sal), saline and proBDNF (Sal-proBDNF), CPP and proBDNF (CPP-proBDNF), NVP and proBDNF (NVP-proBDNF), or Ro25 and proBDNF (Ro25-proBDNF), and then assessed in extinction learning test as above (Fig. 4A). Both CPP

---

**Figure 3.** SD-induced consolidation memory deficit was not attributed to its effect on circadian proBDNF expression. **A,** Conditioned rats were subjected to 4-h SD with or without proBDNF infusion ZT0. At the ZT4 of the following day, to tested fear memory, rats were returned to the same contextual chambers and presented with as few as one tone. **B,** SD impaired memory consolidation of conditioned fear, while injection of cleavage-resistant proBDNF into the vHPC did not rescue the defect; \( ***p < 0.01 \) versus CON; \( n = 7 \) per group. Immunohistochemical detection of proBDNF in vHPC, PFC, striatum, cerebellum, diencephalon and amygdala of SD and control rats. Digital light micrographs (20\( \times \)) of proBDNF-positive cells (**C**) and sum of IOD comparison (**D**); \( \ast p < 0.05 \) versus CON; \( n = 5 \) per group.
To identify whether the proBDNF infusion could effectively reverse the circadian maximum of proBDNF levels, separated groups of rats were chosen and the circadian rhythmicity were detected as above time points. The expression of vHPC proBDNF varies with the circadian cycle (effect of treatment: $F_{(4,35)} = 141.73, p < 0.001$; block 4–6: $p < 0.001$; Fig. 4C) and Ro25 (block 4–6: $p < 0.001$; Fig. 4C) made acquisition of extinction learning more difficulty. However, NVP-proBDNF rats were indistinguishable from Sal-proBDNF rats in their freezing level (block 4: $p < 0.01$, block 5 and 6: $p < 0.001$; Fig. 4C), which demonstrated that proBDNF-NMDA NR2B, but not NR2A, signaling involved in SD-induced deficits in extinction acquisition.

To identify whether the proBDNF infusion could effectively reverse the circadian maximum of proBDNF levels, separated groups of rats were chosen and the circadian rhythmicity were detected as above time points. The expression of vHPC proBDNF varies with the circadian cycle in both CON and SD-proBDNF groups (effect of treatment: $F_{(3,20)} = 57.91, p < 0.001$; Fig. 4D). Both these groups had a higher proBDNF activity at ZT4 than SD group (both comparisons: $p < 0.001$), indicated intrahippocampal infusion of proBDNF maintained the circadian pattern of proBDNF rhythm. However, although injection of mBDNF could slightly enhanced proBDNF level at ZT4 compared SD group ($p < 0.05$), this improvement still lagged behind pro-BDNF group as indicated by the significantly difference between SD-proBDNF and SD-mBDNF at ZT4 ($p < 0.01$). Additionally, we did not observe similar circadian pattern of proBDNF expression in CPP-proBDNF group (data not shown). Collectively these results indicated the infusion of proBDNF were sufficient for maintenance of its circadian rhythm, which could mitigate SD-induced fear extinction deficits.

SD inactivated the proBDNF-mediated vHPC-IL, but not BLA-IL, projection to impair the acquisition of fear extinction

The neural circuits, including vHPC, BLA, and IL-PFC, mediated extinction of conditioned fear (Peters et al., 2010; Sierra-Mercado et al., 2011; Rosas-Vidal et al., 2014). Since HPC and BLA inputs were important for supplying BDNF to the IL-PFC to facilitate extinction memory (Sierra-Mercado et al., 2011; Kimbrough and Biggs, 2014; Rosas-Vidal et al., 2014), we tested if the circadian rhythms of proBDNF activity were co-occurred in three brain regions throughout the circadian cycle (Fig. 5A). Intriguingly, the circadian oscillation was presented in both IL-PFC (compared ZT4 to ZT16: $p < 0.01$; Fig. 5B, left) and vHPC (compared ZT4 to ZT16: $p < 0.01$; Fig. 5B, left), but not BLA, while they disappeared when tested on day following SD (comparison between SD and CON in IL or HPC at ZT4: both $p < 0.01$; Fig. 5B, right). To detect whether the levels of proBDNF in the vHPC affect proBDNF levels in the IL-PFC, anti-proBDNF antibody (Anti) or cleavage-resistant proBDNF (Pro) was injected into the vHPC, and the proBDNF expression of the IL-PFC was assessed 0.5 h following the injection. The infusion anti-proBDNF antibody into vHPC significantly decreased the levels of proBDNF in the IL-PFC (effect of treatment: $F_{(2,18)} = 86.92, p < 0.001$; posthoc: $p < 0.01$; Fig. 5C, left), while the infusion of proBDNF significantly increased IL-PFC proBDNF levels (effect of treatment: $F_{(2,18)} = 136.51, p < 0.001$; posthoc: $p < 0.01$; Fig. 5C, right). Furthermore, the levels in the IL-PFC of the rats (Tra), which were subjected to extinction training, were detected 24 following training. The IL-PFC proBDNF was

**Figure 4.** The mitigating effect of proBDNF on SD-induced deficits in acquisition of fear extinction required NR2B subunit of NMDA receptors. **A,** Treatment procedure. Conditioned rats received post-SD infusion(s) and then tested 48 h following contextual conditioning. **B,** The effect of proBDNF, mBDNF, and anti-proBDNF antibody on SD-induced extinction impairment; ***$p < 0.001$; SD or CON-Anti versus CON or SD-proBDNF; ###$p < 0.001$; SD-mBDNF versus CON or SD-proBDNF; %$p < 0.01$; SD-mBDNF versus SD or CON-Anti; $n = 7$ per group. **C,** Rats were subjected SD between ZT0 and ZT4, the effect of co-infusion NMDA receptor antagonists (CPP, NVP, and Ro25) and proBDNF on freezing levels were tested; ***$p < 0.001$; Sal-proBDNF versus Sal-Sal, CPP-proBDNF or Ro25-proBDNF; ###$p < 0.001$; NVP-proBDNF versus Sal-Sal, CPP-proBDNF or Ro25-proBDNF; $n = 8$ per group. **D,** The proBDNF levels of SD rats that treated with proBDNF and mBDNF throughout the circadian cycle; ***$p < 0.001$; SD versus CON or SD-proBDNF; %$p < 0.01$; SD-mBDNF versus CON or SD-proBDNF; $p < 0.05$; SD-mBDNF versus SD; $n = 6$ per time point.
significantly enhanced compared with Anti and Con groups (both \( p < 0.01 \)), but no statistical difference was found between Tra and Pro groups.

To seek whether maintaining of the co-occurred circadian rhythms was able to rescue the deteriorated effect of SD on extinction acquisition, rats were trained in fear conditioning, rats received post-SD infusion(s) and then tested at the following ZT4. E. The effect of co-infusion of saline into the IL and proBDNF into vHPC [Sal(IL)-Pro(vHPC)], saline into the IL and proBDNF into BLA [Sal(IL)-Pro(BLA)], proBDNF antibody into the IL and saline into vHPC [Anti(IL)-Pro(vHPC)], proBDNF into the IL and saline into vHPC [Pro(IL)-Sal(vHPC)], saline into the IL and saline into vHPC [Sal(IL)-Sal(vHPC)] on freezing level in SD rats; **\( p < 0.01 \): Sal(IL)-Pro(vHPC) or Pro(IL)-Sal(vHPC) versus Sal(IL)-Pro(BLA), Anti(IL)-Pro(vHPC) or Sal(IL)-Sal(vHPC); \( n = 7 \) per group.

**SD suppressed IL-PFC neural activity during extinction of conditioned fear**

Our results clearly indicated that the circadian rhythm of proBDNF in vHPC could influence extinction via IL-PFC. Given the essential role of proBDNF in regulation of neural activity (Nagappan et al., 2009; Yang et al., 2014; Glerup et al., 2016) and the facilitation of IL-PFC excitability during extinction training (Santini et al., 2008; Milad and Quirk, 2012), in a separated group of fear-conditioned rats we sleep deprived preceded by proBDNF infusion or not, and the single-unit activity of IL-PFC was assessed following extinction learning (Fig. 6A). Neuron spike trains were classified by wave form shape (Fig. 6B) and spiking patterns (Fig. 6C). Three hundred sixteen single units were isolated from the IL of nine rats during three recording sessions. As shown in Figure 6B,C, wide-wave form neurons were classified as regular-firing neurons (87 from CON group, 87.9% of group population; 91 from SD group, 87.5% of the population; 97 from SD-proBDNF group, 85.8% of the population) while narrow-wave form neurons were classified as FS interneurons (12 from control group, 12.1% of the population; 13 from SD group, 12.5% of the population; 16 from SD-proBDNF group, 14.2% of the population). Additionally, 17 cells were not classified, since they could not be identified clearly (Fig. 6C). SD group showed a significant increase in basal firing rate of RS projection neurons compared to CON (\( p < 0.01 \)) and SD-proBDNF (\( p < 0.01 \)) group (Fig. 6D, inset). However, the basal firing rate of FS inhibitory interneurons...
was comparable among three groups (Fig. 6E, inset). Following fear extinction training, the normalized spiking frequency of RS neurons from SD group was diminished compared to those from CON group (effect of treatment: $F_{(2,272)} = 268.19, p < 0.001$; block 1–5: $p < 0.001$; Fig. 6D), while infusion of proBDNF resulted in an elevation of firing rate (block 1–5: $p < 0.001$; Fig. 6D), even turn back to the normal level. No main effect of treatment was found in FS interneurons ($F_{(2,38)} = 1.53, p > 0.05$; Fig. 6E). Additionally, Z-scores were also calculated and no statistical difference was found in putative projection cell or inhibitory interneuron (data not shown). These findings indicated that SD-induced circadian proBDNF rhythm disruption may impair fear extinction by suppressing post-training neural activity.

**Discussion**

Endogenous processes referred to as circadian oscillators generate many of the daily rhythms in physiology and behavior of a variety of animals including humans. The present study provides multiple lines of evidence demonstrating that proBDNF activity in the vHPC maintained circadian oscillations, while SD between ZT0 and ZT4 severely lowered the enhancement during the day phase. We discovered SD suppressed acquisition of conditioned fear extinction was associated with the peak of proBDNF, which involved in the function of NR2B subunit of NMDA receptor, during circadian cycle. Furthermore, SD attenuated the projection of vHPC-IL as indicated by the synchronous absent circadian proBDNF in the IL-PFC and the abnormal facilitation of basal firing rate of RS neurons. Infusion of proBDNF to activate the circadian oscillations between vHPC and IL-PFC could effectively mitigate SD-induced deficits in acquisition of extinction and elevate sustained extinction-related activity of RS neurons in the IL-PFC.

Recently, growing researches have been renewed interest in circadian influence in learning and memory (Gerstner and Yin, 2010; Michel et al., 2013). Previous studies pointed on the circadian mBDNF expression and its related signaling were required for memory consolidation and stabilization (Eckel-Mahan et al., 2008; Gerstner and Yin, 2010; Coria-Lucero et al., 2016). Here we found vHPC proBDNF, the precursor of mBDNF, facilitated the extinction of conditioned fear memory, while ablation of proBDNF oscillations by SD, interfering the early stage of sleep or pharmacological blocking proBDNF activity during sleep, resulted in deficits in acquisition of extinction and elevate sustained extinction-related activity of RS neurons in the IL-PFC.
defects even in the absence of early-stage sleep. Several studies showed that proBDNF preferentially activated p75NTR to mediate neuronal cell apoptosis and neurogenesis (Je et al., 2012; Sun et al., 2012), which were required for early stage of sleep to selectively trim newly formed synapses in eliminating and maintaining of memory traces (Griffith and Rosbash, 2008; Vyazovskiy et al., 2008; Li et al., 2017). Compared with robustly effect of proBDNF, we also found that infusion of mBDNF tended to some extent rescue extinction defects. The effect from mBDNF itself or the feedback excitation of mBDNF on proBDNF could be the potential explanations, although the proximate mechanisms were not known. Therefore, in agreement with previous findings (Griffith and Rosbash, 2008; Yang et al., 2014), proBDNF most probably played a critical role in sleep homeostasis to remodel neural circuitry.

Memory traces were typically thought to manifest as plastic changes in neuronal physiology. By activation of NR2B-containing NMDA receptors, proBDNF directly facilitated hippocampal LTD (Yang et al., 2014; Mizui et al., 2015), which served to eliminate fear memory through weakening the unused synaptic connection (Dalton et al., 2012; Li et al., 2016). Fear extinction depended on NMDARs activation (Peters et al., 2010; Dalton et al., 2012). Extendingly, we identified an obligatory role for NR2B-containing NMDA receptors in conveying the effect of circadian proBDNF on acquisition of extinction. It was concurred with previous findings that the function of NR2B-, but notNR2A-, containing NMDA receptors involved in the enhancement of acquisition of extinction memory (Corcoran et al., 2013; Otis et al., 2014). Furthermore, neural correlates of plasticity-related neurophysiology were also regulated by circadian rhythms (Chaudhury et al., 2005; Gerstner and Yin, 2010). Intriguingly, we also found that SD enhanced basal excitability in IL-RS neurons but not inhibitory FS interneurons, which was confirmed by that SD disrupted to establish the tight balance between excitation and inhibition (Griffith and Rosbash, 2008; Vyazovskiy et al., 2009). This excitation-inhibition balance may also contribute to hippocampal proBDNF/p75NTR signaling during subjective night (Riffault et al., 2014). Indeed, the synaptic homeostasis hypothesis of sleep-wake regulation proposed a homeostatic increase in net synaptic strength and neural excitability along with decreased inducibility of associative synaptic potentiation due to saturation after SD (Kuhn et al., 2016). Given that proBDNF/p75NTR mediated synaptic depression boosted network performance by protecting neural networks from saturation (Fino et al., 2005; He et al., 2015), the degradation of circadian proBDNF expression induced by SD most probably affected homeostatic changes in neuronal synchronization. Noteworthy, we demonstrated that the inhibited neural responses that occurred post-acquisition of conditioned extinction was not due to the enhanced neuronal excitability, since the normalized IL-RS frequency from SD was still lowered. In agreement with this observation, cortical mean firing rates changed as a function of sleep homeostasis, and this higher firing sustained in behavioral states (Vyazovskiy et al., 2009). Therefore, the circadian effect on neural excitability could be considered as naturally occurring form of metaplasticity (Finnie and Nader, 2012), in that the synaptic efficacy for an identical stimulation varied based on light/dark cycle (Chaudhury et al., 2005).

In contrast to the well-established function of HPC-BLA-mPFC circuit during contextual fear conditioning (Sotres-Bayon et al., 2009; Sierra-Mercado et al., 2011), we observed that infusing proBDNF into vHPC, but not BLA, attenuated learning extinction deficits in SD condition. Furthermore, blocking IL with anti-proBDNF antibody eliminated the proBDNF effect in the vHPC. Although there was no evidence that proBDNF was directly released from the vHPC inputs to IL-PFC, our findings clearly implied proBDNF infused into vHPC could affect the expression of proBDNF in the IL-PFC. The possible explanation is the infusion of proBDNF could be directly or indirectly transported into IL or the infusion of proBDNF could increase vHPC excitability leading to increased release of proBDNF in IL. Further evidence may come from transgenic animals and optogenetics. Several findings pointed out SD induced the aberrant HPC-PFC communication, as others have hypothesized that similar findings were a result of decreased consolidation of HPC-based memories during sleep (Mander et al., 2013) or improper HPC engagement during PFC-mediated tasks (Egan et al., 2003). For example, an increase in HPC activity and reduced long-term memory recall following sleep was observed with decreased synchronized activity between HPC and PFC (Mander et al., 2013). Other lab reported an exaggerated increase in low theta power in the HPC of Bdnf-e4 mice during freezing (Hill et al., 2016). This observation was consistent with HPC over-activation in individuals with the Val66Met polymorphism (Mascetti et al., 2013). Taken together, our results supported the vHPC-IL, but not BLA-IL, circuit engaged in proBDNF-mediated sleep homeostasis.

The combined rodent-human extinction findings predicted that extinguish fear responses and its associated network would be impaired in post-traumatic stress disorder (PTSD) patients (Milad et al., 2009; Sierra-Mercado et al., 2011). Consistent with mPFC dysfunction, PTSD patients showed normal conditioning and within-session extinction, but were unable to recall extinction memory (Milad et al., 2009). In healthy subjects, acquisition and retrieval of extinction was associated with increased activity in the vmPFC (homologous to rodent IL) and HPC (Milad et al., 2007). PTSD subjects exhibited deactivation in these areas, coupled with hyperactivation of the dorsal anterior cingulate cortex (homologous to rodent PL; Milad et al., 2009). Our findings suggested that it was during extinction training when activity in these areas was important. Sleep was maintained or disturbed might predict an individual’s ability to recover from a preexisting psychiatric condition (Pace-Schott et al., 2015; Kuhn et al., 2016; Schmitt et al., 2016). Restorative sleep was necessary to maintain adequate BDNF activity, which was notably reduced in patients with PTSD and was increased following psychiatric treatment (Felmingham et al., 2013; Zhang et al., 2014). Therefore, sleep was a key mediator at the
connection between trauma exposure and the BDNF system (Andero and Ressler, 2012; Milad and Quirk, 2012), for which deregulation can exacerbate the mental and physical comorbidity (Andero and Ressler, 2012; Hill et al., 2016). Our study given the prospect that improved sleep quality may result in psychiatric recovery, which was associated with reactivity proBDNF signaling.

In conclusion, the proBDNF expression underwent circadian oscillations in HPC-IL pathway that correlated with the ability to acquisition of conditioned extinction. SD induced deficits in extinction learning, but not conditioning acquisition. The disruption of the circadian proBDNF levels induced by SD involved in NR2B, but not NR2A, subunits of NMDA receptors. Furthermore, SD enhanced the basal firing rate but depressed response activity of IL-RS neurons, which was reversed by providing proBDNF during early stage of sleep. Our findings suggested that proBDNF quite possible engaged in regulation of sleep homeostasis by balancing excitation and inhibition in the projection of HPC-IL. Further studies should focus on the role of proBDNF activity in extinguishing unwanted memory to offer potential therapeutic strategies for psychiatric disorders.

References

Ackermann S, Rasch B (2014) Differential effects of non-REM and REM sleep on memory consolidation? Curr Neurol Neurosci Rep 14:430.

An L, Sun W (2018) Acute melamine affects spatial cognition and hippocampal synaptic plasticity by presynaptic and postsynaptic inhibition of glutamatergic transmission in adolescent offspring. Toxicol Lett 269:55–64.

An L, Li X, Tang C, Xu N, Sun W (2018) Hippocampal proBDNF facilitates place learning strategy associated with neural activity in rats. Brain Struct Funct 223:4099–4113.

Andero R, Ressler KJ (2012) Fear extinction and BDNF: translating animal models of PTSD to the clinic. Genes Brain Behav 11:503–511.

Bai YY, Ruan CS, Yang CR, Li JY, Kang ZL, Zhou L, Liu D, Zeng YQ, Wang TH, Tian CF, Liao H, Bobrovskaya L, Zhou XF (2016) ProBDNF signaling regulates depression-like behaviors in rodents under chronic stress. Neuropsychopharmacology 41:2882–2892.

Bekinschtein P, Cammarota M, Katche C, Slipczuk L, Rossato JI, Goldin A, Izquierdo I, Medina JH (2008) BDNF is essential to promote persistence of long-term memory storage. Proc Natl Acad Sci USA 105:2711–2716.

Blanchard RJ, Blanchard DC (1969) Crouching as an index of fear. J Comp Physiol Psychol 67:370–375.

Boyce R, Glasgow SD, Williams S, Adamantidis A (2016) Causal evidence for the role of REM sleep theta rhythm in contextual memory consolidation. Science 352:812–816.

Burgos-Robles A, Vidal-Gonzalez I, Santini E, Quirk GJ (2007) Consolidation of fear extinction requires NMDA receptor-dependent LTD in rats. Neuroscience 11:1074–1082.

Egan MF, Koijima M, Callicott JH, Goldberg TE, Kolachana BS, Bertolino A, Zaitsev E, Gold B, Goldman D, Dean M, Lu B, Weinberger DR (2003) The BDNF val66met polymorphism affects activity-dependent secretion of BDNF and human memory and hippocampal function. Cell 112:257–269.

Fan YJ, Wu LL, Li HY, Wang YJ, Zhou XF (2008) Differential effects of pro-BDNF on sensory neurons after sciatic nerve transection in neonatal rats. Eur J Neurosci 27:2380–2390.

Flemingham KL, Dobson-Stone C, Schofield PR, Quirk GJ, Bryant RA (2013) The brain-derived neurotrophic factor Val66Met polymorphism predicts response to exposure therapy in posttraumatic stress disorder. Biol Psychiatry 73:1059–1063.

Finnie PS, Nader K (2012) The role of metaplasticity mechanisms in regulating memory destabilization and reconsolidation. Neurosci Biobehav Rev 36:1667–1707.

Fino E, Glowinski J, Venance L (2005) Bidirectional activity-dependent plasticity at corticostriatal synapses. J Neurosci 25:11279–11287.

Gerstner JR, Yin JC (2010) Circadian rhythms and memory formation. Nat Rev Neurosci 11:577–588.

Glerup S, Bolcho U, Mølgaard S, Bøggild S, Vaegter CB, Smith AH, Gentile F, Haugebo P, Van der Zee EA, Meerlo P (2010) Coping with sleep deprivation: shifts in regional brain activity and learning strategy. Sleep 33:1465–1473.

He K, Huertas M, Hong SZ, Tye X, Hell JW, Shouval H, Kirkwood A (2015) Distinct eligibility traces for LTP and LTD in cortical synapses. Neuron 88:528–538.
Hill JL, Hardy NF, Jimenez DV, Maynard KR, Kardian AS, Pollocock J, Schloesser RJ, Martinowich K (2016) Loss of promoter IV-driven BDNF expression impacts oscillatory activity during sleep, sensory information processing and fear regulation. Transl Psychiatry 6:e873.

Holmes M, Nieto-Gonzalez JL, Vardy I, Vaegter CB, Nykaeyer A, Jensen K (2009) Mature BDNF, but not proBDNF, reduces excitability of fast-spiking interneurons in mouse dentate gyrus. J Neurosci 29:12412–12418.

Insel N, Barnes CA (2015) Differential activation of fast-spiking and regular-firing neuron populations during movement and reward in the dorsal medial frontal cortex. Cereb Cortex 25:2631–2647.

Itoh N, Enomoto A, Nagai T, Takahashi M, Yamada K (2016) Molecular mechanism linking BDNF/TrkB signaling with the NMDA receptor in memory: the role of Girdin in the CNS. Rev Neurosci 27:481–490.

Jang SW, Liu X, Pradoldej S, Tosini G, Chang Q, Iuvone PM, Ye K (2017) N-acetylsperotonin activates TrkB receptor in a circadian rhythm. Proc Natl Acad Sci USA 107:3876–3881.

Je HS, Yang F, Ji Y, Nagappan G, Hempstead BL, Lu B (2012) Role of pro-brain-derived neurotrophic factor (proBDNF) to mature BDNF conversion in activity-dependent competition at developing neuromuscular synapses. Proc Natl Acad Sci USA 109:15924–15929.

Kim H, Åhrlund-Richter S, Wang X, Deisseroth K, Carlén M (2016) Prefrontal parvalbumin neurons in control of attention. Cell 164: 208–218.

Kimbrough A, Biggs LM (2014) BDNF signaling potentiates transmission of information from the basolateral amygdala to infralimbic prefrontal cortex during conditioned taste aversion extinction. J Neurosci 34:12617–12618.

Kopp C, Longordo F, Nicholson JR, Lüthi A (2006) Insufficient sleep reversibly alters bidirectional synaptic plasticity and NMDA receptor function. J Neurosci 26:12456–12465.

Kuhn M, Wolf E, Maier JG, Mainberger F, Schmid H, Bürklein M, Kojima M (2015) BDNF pro-peptide actions facilitate hippocampal LTD and are altered by the common BDNF polymorphism Val66Met. Proc Natl Acad Sci USA 112:E3067–E3074.

Mueller D, Bravo-Rivera C, Quirk GJ (2010) Infralimbic D2 receptors are necessary for fear extinction and extinction-related tone responses. Biol Psychiatry 68:1055–1060.

Nagappan G, Zaitsev E, Senatorov V Jr, Yang J, Hempstead BL, Lu B (2009) Control of extracellular cleavage of proBDNF by high frequency neuronal activity. Proc Natl Acad Sci USA 106:1267–1272.

Nakai T, Nagai T, Tanaka M, Itoh N, Asai N, Enomoto A, Asai M, Yamada S, Saifullah AB, Sokabe M, Takahashi M, Yamada K (2014) Girdin phosphorylation is crucial for synaptic plasticity and memory: a potential role in the interaction of BDNF/TrkB/Akt signaling with NMDA receptor. J Neurosci 34:14995–15008.

Ots J, Gómez-Pinilla F, Müller D (2014) Infralimbic BDNF/TrkB enhancement of GluN2B currents facilitates extinction of a cocaine-conditioned place preference. J Neurosci 34:6057–6064.

Oyanedel CN, Kelemen E, Scheller J, Born J, Rose-John S (2015) Peripheral and central blockade of interleukin-6 trans-signaling differentially affects sleep architecture. Brain Behav Immun 50:178–185.

Pace-Schott EF, Germain A, Milad MR (2015) Effects of sleep on memory for conditioned fear and fear extinction. Psychol Bull 141:835–857.

Paterson NE, Froestl W, Markou A (2005) Repeated administration of the GABAB receptor agonist CGP44532 decreased nicotine self-administration, and acute administration decreased cue-induced reinstatement of nicotine-seeking in rats. Neuropsychopharmacology 30:119–128.

Paxinos G, Watson C (1997) The rat brain in stereotaxic coordinates (third edition). New York: Academic Press.

Peters J, Diepma-Perea LM, Melendez LM, Quirk GJ (2010) Induction of fear extinction with hippocampal-infralimbic BDNF. Science 328:1288–1290.

Rasch B, Born J (2013) About sleep’s role in memory. Physiol Rev 93:681–766.

Rasch B, Büchel C, Gais S, Born J (2007) Odor cues during slow-wave sleep prompt declarative memory consolidation. Science 315:1426–1429.

Ravassard P, Hamieh AM, Joseph MA, Fraize N, Libourel PA, Leparc G, Arthaud L, Meisslir C, Touret M, Mellaret G, Salin PA (2016) REM sleep-dependent bidirectional regulation of hippocampal-based emotional memory and LTP. Cereb Cortex 26:1488–1500.

Riffault B, Medina I, Dumen C, Thalman C, Ferrand N, Fiedel P, Gairars JA, Porcher C (2014) Pro-brain-derived neurotrophic factor inhibits GABAergic neurotransmission by activating endocytosis and repression of GABAA receptors. J Neurosci 34:13516–13534.
Rolls A, Makam M, Kroeger D, Colas D, de Lecea L, Heller HC (2013) Sleep to forget: interference of fear memories during sleep. Mol Psychiatry 18:1166–1170.

Rosas-Vidal LE, Do-Monte FH, Sotres-Bayon F, Quirk GJ (2014) Hippocampal-prefrontal BDNF and memory for fear extinction. Neuropsychopharmacology 39:2161–2169.

Santini E, Quirk GJ, Porter JT (2008) Fear conditioning and extinction differentially modify the intrinsic excitability of infralimbic neurons. J Neurosci 28:4028–4036.

Schmitt K, Holsboer-Trachsler E, Eckert A (2016) BDNF in sleep, insomnia, and sleep deprivation. Ann Med 48:42–51.

Sierra-Mercado D, Padilla-Coreano N, Quirk GJ (2011) Dissociable roles of prelimbic and infralimbic cortices, ventral hippocampus, and basolateral amygdala in the expression and extinction of conditioned fear. Neuropsychopharmacology 36:529–538.

Sotres-Bayon F, Diaz-Mataix L, Bush DE, LeDoux JE (2009) Dissociable roles for the ventromedial prefrontal cortex and amygdala in fear extinction: NR2B contribution. Cereb Cortex 19:474–482.

Stickgold R, Walker MP (2013) Sleep-dependent memory triage: evolving generalization through selective processing. Nat Neurosci 16:139–145.

Sun W, Li X, An L (2018a) Distinct roles of prelimbic and infralimbic proBDNF in extinction of conditioned fear. Neuropsychopharmacology 131:11–19.

Sun W, Li X, Tang C, An L (2018b) Acute low alcohol disrupts hippocampus-prefrontal neural correlate of learning strategy by inhibition of PKA/CREB pathway in rats. Front Pharmacol 9:1439.

Sun Y, Lim Y, Li F, Liu S, Lu JJ, Haberberger R, Zhong JH, Zhou XF (2012) ProBDNF collapses neurite outgrowth of primary neurons by activating RhoA. PLoS One 7:e35883.

Tadavarty R, Kaan TK, Sastry BR (2009) Long-term depression of excitatory synaptic transmission in rat hippocampal CA1 neurons following sleep-deprivation. Exp Neurol 216:239–242.

Vianna MR, Szapiro G, McGaugh JL, Medina JH, Izquierdo I (2001) Retrieval of memory for fear-motivated training initiates extinction requiring protein synthesis in the rat hippocampus. Proc Natl Acad Sci USA 98:12251–12254.

Vyazovskiy VV, Cirelli C, Pfister-Genskow M, Faraguna U, Tononi G (2008) Molecular and electrophysiological evidence for net synaptic potentiation in wake and depression in sleep. Nat Neurosci 11:200–208.

Vyazovskiy VV, Olcese U, Lazimy YM, Faraguna U, Esser SK, Williams JC, Cirelli C, Tononi G (2009) Cortical firing and sleep homeostasis. Neuron 63:865–878.

Vyazovskiy VV, Palchykova S, Achermann P, Tobler I, Deboer T (2017) Different effects of sleep deprivation and torpor on EEG slow-wave characteristics in Djungarian hamsters. Cereb Cortex 27:950–961.

Walker AG, Wenthr CJ, Xiang Z, Rook JM, Emmite KA, Niswender CM, Lindsley CW, Conn PJ (2015) Metabotropic glutamate receptor 3 activation is required for long-term depression in medial prefrontal cortex and fear extinction. Proc Natl Acad Sci USA 112:1196–1201.

Wiltgen BJ, Sanders MJ, Anagnostaras SG, Sage JR, Fanselow MS (2006) Context fear learning in the absence of the hippocampus. J Neurosci 26:4028–4036.