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Research report

Photoperiod alters fear responses and basolateral amygdala neuronal spine density in white-footed mice (Peromyscus leucopus)


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HIGHLIGHTS

- Short days lengthen enhance fear memory in white-footed mice (Peromyscus leucopus).
- Short days increase spine density in the basolateral amygdala.
- Short days do not alter dendritic spines or arborization in the infralimbic PFC.

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ABSTRACT

Photoperiodism is a biological phenomenon in which environmental day length is monitored to ascertain time of year to engage in seasonally appropriate adaptations of physiology and behavior. These seasonal changes in physiology and behavior can be induced in a laboratory setting by simply exposing animals to different static day lengths [1]. White-footed mice (Peromyscus leucopus) display a suite of changes in physiology and behavior induced by exposure to short days, including changes in behavior, brain volume and functional connectivity, and enhanced HPA axis reactivity [2–4]. In recent years, photoperiodic rodent models are finding more utility as models of human pathologies involving alterations in brain structure and function [5,6].

Among human post-traumatic stress disorder (PTSD) patients, reduced hippocampal volume is associated with susceptibility to PTSD, but not severity of symptoms [7], amygdala activity is associated with symptom severity [8], and PTSD patients have enhanced HPA axis negative feedback [7]. Additionally, the medial prefrontal cortex (mPFC) is hyporesponsive to fear-related stimuli in patients suffering from PTSD [9,10]. Translational rodent studies have identified these three brain regions as critical components of the neural circuit underlying associative fear memory. The basolateral amygdala (BLA), hippocampus, and mPFC all play critical roles in fear memory: complex modulation of reciprocal connections among the mPFC, hippocampus, and basolateral amygdala (BLA) are integral to associative fear memory [7,11–13].

Although standard laboratory rodents (Rattus norvegicus and Mus musculus) have been widely used to model PTSD and other human psychiatric disorders, to fully understand combined...
factors underlying the development, maintenance, and treatment of these diseases, more diverse animal models are needed [5]. Toward this end, photoperiod-induced changes in the brain of white-footed mice mirror several components of the etiology of PTSD. For example, in common with the human PTSD condition, short-day exposed white-footed mice have reduced hippocampal volume [4] and increased HPA axis feedback [3]. To our knowledge, photoperiod-mediated changes in fear memory and responses in this, and other, photoperiodic rodent species remain largely undescribed. A preliminary study in our lab indicated that, unlike short-day induced impairments in hippocampal-mediated spatial learning and memory, white-footed mice exposed to short days may have enhanced non-spatial fear memory in the passive avoidance test [4].

Based on the commonality of photoperiodic changes in white-footed mice with the etiology of PTSD described above, and on preliminary data demonstrating short day enhancement of fear memory, we hypothesized that exposure to short days would enhance fear memory and alter neuronal morphology in brain regions implicated in associative fear memory. To test our hypothesis, we exposed male white-footed mice to either short or long day lengths for ten weeks to induce maximal photoperiodic responses, tested them in an auditory fear conditioning test, and examined the neuronal morphology in the BLA and the infralimbic region of the mPFC (IL) using Golgi-Cox staining.

2. Materials and methods

2.1. Animals

Nineteen adult (±65 d of age) male P. leucopus, from our breeding colony maintained at the Ohio State University, were randomly assigned to either long (LD: 16L:8D, n = 10) or short day lengths (SD: 8L:16D, n = 9) for 10 weeks to establish photoperiod-induced changes prior to behavioral testing [2,4]. Mice were housed in standard polycarbonate cages (32 cm × 18 cm × 14 cm), maintained at constant temperature (21 ± 4 °C) and relative humidity (50 ± 5%), provided ad libitum access to filtered tap water and food (Harlan Teklad 8640, Indianapolis, IN, USA), and received care from the Ohio State University Laboratory Animal Resource staff for the duration of the study. All procedures proposed were approved by the Ohio State University Institutional Animal Care and Use Committee and are in compliance with guidelines established by the National Institutes of Health [14].

2.2. Behavioral test

2.2.1. Auditory fear conditioning

To assess tone-conditioned fear acquisition and retention, 19 (n = 10 LD; n = 9 SD) mice were assessed using the Near-IR Video Fear Conditioning System (Med Associates Inc., St. Albans, VT, USA). For each animal, the tone-conditioned fear during the light phase mice were placed directly from their vivarium rooms and were placed in the test chamber illuminated with white light for a 2 min habituation period with 68 dB white noise. Mice were then exposed to a series of 8 conditional stimuli (80 dB tone, CS) for 6 s with the last 2 s paired with a 0.75 mA foot shock (unconditioned stimulus, US). Mice remained in the chamber for an additional 60 s after the last CS/US pairing before being returned to their home cages. Freezing behavior was recorded by the software for the 2 min baseline, during the first 4 s of each tone, during the 30 s interval between CS presentations, and for the 60 s after the final CS. To access contextual fear retention, 24 h after the acquisition session, mice were placed in the original unmodified chamber and freezing behavior to the unmodified chamber was recorded for 60 s, and mice were returned to their home cages. Four hours after the contextual fear retention test, mice were tested for retention of the CS–US pairing with the following modifications to alter context. Mice were transported from their vivarium rooms via sound- and light-attenuating boxes to a staging area. From there, mice were brought into the testing room lit with dim red light and placed into the chambers. To avoid context-dependent freezing, the chamber was modified via the addition of a smooth plastic floor, a semi-circular retinal testing chamber, lights were extinguished, and a gauze pad with a drop of vanilla extract was placed in the chamber to present the CS in a novel environment. Mice were then tested for retention of the CS–US pairing by using the protocol described above for the acquisition trial above without receiving the foot shock (US). Zoological studies of auditory fear conditioning, under deep isoflurane anesthesia, mice were exsanguinated via the retro-orbital sinus, plasma was collected from the blood samples as previously described [4], and samples were stored at −80°C for corticosterone assay. Immediately after blood collection, mice were rapidly decapitated and brains were processed to study neuronal morphology (after [4,15]) using a commercially available Golgi–Cox impregnation kit (FD NeuroTechnologies, Ellicott City, MD, USA) according to the manufacturer's instructions. Briefly, after impregnation, brains were cut into 100 μm coronal sections and thaw mounted on to gelatin-coated slides. Slides were then developed, counterstained with cresyl violet acetate, dehydrated with xylenes, and coverslipped with Permount (Fisher).

2.2.2. Photoperiodicity

Mice were exposed to either short (SD: 8L:16D) or long day lengths (LD: 16L:8D) for 10 weeks to induce maximal photoperiodic responses, tested them in an auditory fear conditioning test, and examined the neuronal morphology in the BLA and the infralimbic region of the mPFC (IL) using Golgi-Cox staining.

2.3. Sample collection and histology

Twenty hours after the completion of auditory fear conditioning, under deep isoflurane anesthesia, mice were exsanguinated via the retro-orbital sinus, plasma was collected from the blood samples as previously described [4], and samples were stored at −80°C for corticosterone assay. Immediately after blood collection, mice were rapidly decapitated and brains were processed to study neuronal morphology (after [4,15]) using a commercially available Golgi–Cox impregnation kit (FD NeuroTechnologies, Ellicott City, MD, USA) according to the manufacturer's instructions. Briefly, after impregnation, brains were cut into 100 μm coronal sections and thaw mounted on to gelatin-coated slides. Slides were then developed, counterstained with cresyl violet acetate, dehydrated with xylenes, and coverslipped with Permount (Fisher).

2.3.1. Dendritic Arborization

Pyramidal neurons (n = 4–6 for each mouse) in the infralimbic cortex (IL), identified by its cytoarchitecture and neuroanatomical position medial to the forepaw minor and cingulum between 1.3 and 1.9 mm anterior to bregma [16], were traced at 400 × and quantified using neuronal tracing software (Neurolucida, MicroBrightfield, VT, USA). Neurons were traced only if they met the following criteria: (1) completely and uniformly impregnated with Golgi stain, (2) all dendrites were intact and visible, and (3) not obscured by other stained neurons (after [4]). The basolateral amygdala (BLA), identified by its location bounded by the branchial arms of the external capsule between 0.8 and 2.0 mm posterior to bregma [16], did not contain sufficient numbers of neurons that met the above criteria for analysis. Representative values for each parameter measured by the software (see section 3) from each animal were calculated by averaging values from all neurons traced. Representative values calculated for each animal were then used for further analysis.

2.3.2. Dendritic Spine Density Analysis

Dendritic spines of the neurons were traced at 1000× using NeuroLucida software (Microbrightfield, VT, USA). Within the BLA, for each animal average spine density was calculated by selecting six neurons, and an unbranched, unbroken, and consistently stained dendritic segment at least 50 μm away from the soma from each neuron was quantified. Any protrusion originating from the dendritic shaft was classified as a spine, and all spines along a continuous 80 μm segment were counted for spine density analysis (after [17]). For the IL, average spine densities were quantified for each animal from both basal and apical dendrites, selected as above. For each neuron a total of 80 μm of basilar and 80 μm apical dendrite were quantified for spine density. For each brain region, the average spine density calculated from each animal was then used for further comparative analysis.

2.3.3. Corticosterone assay

Frozen plasma samples were thawed on ice and assayed for corticosterone using a commercially available double antibody RIA kit according to manufacturer's instructions (Cat# 07120102; MP Biomedical, Costa Mesa, CA, USA). The intra-assay coefficient of variation was 7.9%.

2.3.4. Statistics

Repeated measures ANOVA were used to compare auditory fear responses over time and dendritic arborization measures (Sholl analysis). Student’s t tests were used for comparisons between photoperiods for physiological data, spine density analysis, and for follow up testing of specific time points within trials for auditory fear testing after a main effect was identified by ANOVA. Data with unequal variance were log transformed prior to analysis. All analyses were performed using SPSS software (v19; IBM, NY, USA) and differences were considered statistically significant at p < 0.05.

3. Results

3.1. Physiological measures

Exposure to short day lengths did not affect body mass (t17=0.185, p > 0.05) or inguinal fat pad mass (t17=0.036, p > 0.05). Compared to LD-exposed counterparts, exposure to SD reduced masses of all reproductive tissues assessed (paired ttests, t15=4.934, p < 0.001; epididymides, t17=2.872, p < 0.05; seminal vesicles, t17=2.655, p < 0.05 Fig. 1, left). Short day exposure did not affect basal corticosterone concentrations at the terminal bleed (t16=0.904, p > 0.05; Fig. 1, right).

3.2. Behavioral measures

3.2.1. Auditory-cued Fear Conditioning

LD and SD mice did not differ in their freezing responses during acquisition of the CS–US pairing across trials (repeated measures ANOVA: F(1,15) = 0.056, p > 0.05: Fig. 2A). Comparing baseline to post CS–US presentation, both SD and LD mice increased freezing behavior across acquisition (LD, t8=−2.714, p < 0.05; SD t7=−3.457.
p < 0.05), however photoperiod did not affect increases in freezing between pre and post-stimulus (Fig. 2A). There were no differences due to photoperiod in freezing to context 24 h after acquisition (t\(_{17} = 0.777, p > 0.05\); not shown). Because of individual variance in inter-trial interval freezing behavior, freezing responses during retention tone trials were corrected by subtracting freezing during the 30 s immediately prior to tone presentation from freezing during the tone. Compared to mice exposed to LD, SD exposure increased freezing to tone across retention trials (repeated measures ANOVA: F\(_{1,17} = 8.160, p < 0.05\)). Follow up, within-trial, 3.3. Neuronal morphology

3.3.1. Dendritic spine density

Compared to LD mice, mice exposed to SD had increased spine density on dendrites in the BLA (t\(_{10} = -4.196, p < 0.01\); Fig. 3). In the IL, there were no differences in spine density due to photoperiod in either basilar (t\(_{13} = -0.332, p > 0.05\)) or apical dendrites (t\(_{13} = 0.753, p < 0.05\); Fig. 4A).

3.3.2. IL pyramidal neuronal morphology

No differences due to photoperiod were found in IL pyramidal neurons for cell body perimeter (t\(_{14} = 1.332, p > 0.05\)), cell body area (t\(_{14} = 1.190, p > 0.05\), basilar dendrite length (t\(_{14} = -0.066, p > 0.05\)), apical dendrite length (t\(_{14} = -0.061, p > 0.05\)), or total dendrite length (t\(_{14} = -0.074, p > 0.05\); data not shown). No differences were observed due to photoperiod in number of intersections (repeated measures ANOVA: F\(_{1,14} = 410.144, p > 0.05\); Fig. 4C) or dendrite length (repeated measures ANOVA: F\(_{1,14} = 761.671, p > 0.05\); not shown) via Sholl analysis of dendritic arborization complexity.

4. Discussion

The present study in white-footed mice demonstrates three novel findings. (1) Exposure to short day lengths enhances associative fear memory in the auditory-cued fear conditioning. (2) Photoperiod and fear conditioning interact to increase spine density of the neurons in the BLA. However, (3) neither photoperiod nor the interaction of photoperiod and auditory-cued fear conditioning alter the morphology of pyramidal neurons within the IL. Exposure to short days in white-footed mice reduces hippocampal volume, alters hippocampal dendritic spine density, and impairs LTP within the hippocampus [2,4]. Pursuant to these day length induced morphological and physiological changes, SD mice have impaired hippocampal-dependent spatial learning and memory [2,4,15]. Non-hippocampal learning and memory, assessed by several behavioral tests, was previously reported to be unaffected by photoperiod [4]. However, in the passive avoidance fear memory test as previously demonstrated [4], although SD mice did
Effects of SD exposure and fear conditioning on pyramidal neuron morphology in the infralimbic cortex. (A) SD exposure did not alter spine density on either apical or basilar dendrites. (B) 200× photomicrograph of a representative IL pyramidal neuron (upper) and its Neurolucida reconstruction upon which Sholl analysis was performed (lower). (C) Exposure to SD did not alter complexity of pyramidal cell dendritic arborization (Sholl analysis) of infralimbic cortical pyramidal neurons.

not differ from their LD counterparts in latency to step through during training or testing phases, only SD mice increased their latency to step through after training, indicating there may be photoperiodic alterations in fear-related memory. We have extended these findings to show SD-induced enhancement of fear memory in the auditory-cued fear test (Fig. 2B); supporting our hypothesis that SD exposure enhances fear memory. Additionally, increased spine density in the dendrites of the BLA neurons is associated with increased fear memory in SD mice (Fig. 3). The BLA is critical for encoding fear memory and imparting emotional valence on memories [13,17–19]. Furthermore, enhanced spine density within the BLA is associated with enhanced auditory fear memory [20]. Thus, SD exposure causes hippocampal atrophy and functional impairments in the hippocampus, whereas the amygdala becomes hypertrophic and functionally enhanced by exposure to short day lengths. Although time-of-day differences have been reported in fear learning [21–23], the differences are generally confined to contextual fear, rather than tone-cued fear [21,22]. The current findings are not likely due to differences in circadian patterns of fear learning and memory between SD and LD mice as we conducted our acquisition trials during the light phase when fear learning is strongest [21,22], and all of our testing occurred at the same circadian time for both groups, thus limiting any circadian effects.

What is the role of the IL area of the mPFC in photoperiodic alteration in fear memory? The IL/mPFC is critical for orchestrating the balance of the BLA and the hippocampus in the formation and extinction of emotionally charged memories, such as fear conditioning. Reciprocal connections between the mPFC and the BLA regulate encoding and extinction of fear memories [12]. The mPFC-hippocampus pathway also coordinates extinction of fear memories [20,24,25], and the BLA-hippocampus pathway is critical for modulating fear memory encoding and recall [26–28]. Well-orchestrated coordination of all the three brain regions is critical for encoding, recall, and extinction of fear memories [11,13,29]. Although SD mice do show cognitive inflexibility in spatial reversal learning [4], presumably mediated by the IL [30], the current neuroanatomical findings (Fig. 4) do not support a specific role for altered IL function in the SD enhancement of fear memory. However, the role of photoperiodic impairment in the mPFC-hippocampus pathway in cognitive flexibility in both spatial and fear learning remains to be described.

One of the hallmarks of photoperiodic responses in rodents is the involution of the gonads by exposure to short day lengths (Fig. 1) via inhibition of the hypothalamic-pituitary-gonadal (HPG) axis, which results in low gonadal sex steroid concentrations (reviewed in [1]). Gonadal sex steroids can influence neuronal morphology in the regions above implicated in fear memory [31,32]. Although gonadectomy can recapitulate some of the appropriate photoperiodic responses, gonadal steroids are not the sole contributing factor to the suite of behavioral and physiological responses in photoperiodic rodents (reviewed in [1]). Indeed, gonadal steroids enhance fear memory; estrogens facilitate fear conditioning by upregulating CRH expression in the amygdala [33], and androgens (testosterone and dihydrotestosterone) also facilitate conditioned fear, but the enhanced effects are mediated in the hippocampus, and not the amygdala [34]. Thus, reduced sex steroid concentrations in SD mice should lead to impaired fear memory, which argues strongly against a significant role of gonadal steroids in SD enhancement of fear memory. However, we currently have ongoing experiments to study the effects of sex steroids on hippocampal- and amygdala-mediated memory.

In addition to inhibiting the HPG axis, short days enhance HPA axis responsiveness and negative feedback in white-footed mice [3]. Although photoperiod does not affect baseline corticosterone concentrations (Fig. 1; [3]), SD mice display increased HPA axis responsiveness to stressors and increased negative HPA axis feedback, potentially regulated at the level of the hippocampus by SD elevation of hippocampal glucocorticoid and mineralocorticoid receptors [3]. Photoperiod alterations in the HPG and HPA axis have adaptive significance for this species. To survive the energetic bottleneck of reduced food availability and increased thermogenic demands during the short days of winter, energy is conserved by reduction of reproductive tissue mass and reproduction-associated behaviors (reviewed in [1]). Additionally, activation of the HPG axis (glucocorticoid response to stressors) to mobilize energy for the fight-or-flight response is energetically expensive, thus increased regulation (efficiency) of the HPA axis in short days may be an adaptive response to conserve energy [3].
In the North American temperate zone home range of white-footed mice, photoperiodic changes in habitat alter behavior and distribution [35]. Short day mice inhabit an environment that is both energetically demanding and devoid of dense understory cover, thus the alterations in endocrine responses to stress (HPA axis) and behavioral responses to fearful stimuli (fear memory) described above may provide an adaptive advantage to survive in their winter habitat. During the short days of winter forest understory is greatly reduced and leaf cover is minimal [36], and small photoperiodic mammals, including white-footed mice, alter their habitat use and movement to areas within their home range with maximum leaf understory density for predator avoidance [37–40]. To our knowledge, the effect of photoperiod on predator avoidance has never been directly tested on white-footed mice, however short days increase predator avoidance behavior in other rodent species [41,42]. In short days, enhanced fear memory may be advantageous for survival by offsetting the interaction of reduced available ground cover for predator avoidance with impaired hippocampal-mediated spatial navigation [2,4]. Thus it is possible that the photoperiodic differences in fear responses reported here are ecologically relevant and are an integral part of a suite of adaptive responses to short days, which include changes in physiology and hippocampal function.

In summary, short day exposed white-footed mice have reduced hippocampal volume [4], increased HPA axis negative feedback [3], increased fear memory (Fig. 2), and increased connectivity within the BLA (Fig. 3). Among human PTSD patients, reduced hippocampal volume is associated with susceptibility to PTSD, but not severity of symptoms [7], amygdala activity is positively correlated with symptom severity [8], and PTSD patients have enhanced HPA axis negative feedback [7]. One of the greatest research challenges of mechanisms involved in psychiatric disorders is the difficulty in replicating the symptoms of the disease in animal models [43], and it is important to develop new ethologically relevant models for translational research of psychiatric disorders [5,44]. In the laboratory setting, a simple day length manipulation in white-footed mice, exposure to short days, concurrently replicates three main features of PTSD and enhances fear memory responses. Taken together, the photoperiodic modulation of brain fear circuits and memory in white-footed mice described in this, and previous studies, may argue for the potential utility of this species as an additional, yet unique, animal model to research how a single environmental factor (day length) can interact with genes to alter phenotype to resemble a human psychiatric disorder, such as PTSD.

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References


