Short Photoperiods Impair Spatial Learning and Alter Hippocampal Dendritic Morphology in Adult Male White-Footed Mice (Peromyscus leucopus)

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Although seasonal changes in brain morphology and function are well established in songbirds, seasonal plasticity of brain structure and function remain less well documented in mammals. Nontropical animals display many adaptations to reduce energy use to survive winter, including cessation of reproductive activities. Because of the high energetic costs of brain tissue, we hypothesized that male white-footed mice (Peromyscus leucopus) would reduce brain size in response to short days as well as regress their reproductive systems. Because short days may decrease hippocampal volume and impair spatial learning and memory in rodents and because of the potential for seasonal plasticity in the hippocampus, we hypothesized that photoperiod alters hippocampal morphology to affect spatial learning and memory. Mice housed in either long or short days for 10 weeks were examined for performance in a water maze; brains were then removed and weighed, and hippocampal volumes were determined. We also measured dendritic morphology and spine density in the CA1, CA3, and dentate gyrus. Short days decreased brain mass and hippocampal volume compared with long days. Short days also impaired long-term spatial learning and memory relative to long days but did not affect sensory discrimination or other types of memory. Short days decreased apical (stratum lacunosum-moleculare) CA1 spine density, as well as increased basilar (stratum oriens) CA3 spine density. Results from this study suggest that photoperiod alters brain size and morphology, as well as cognitive function. Understanding the mechanisms mediating these photoperiod-induced alterations may provide insight for treatment of seasonal cognitive and affective disorders.

Key words: seasonality; photoperiod; spatial learning and memory; hippocampus; spine density; dendritic complexity

Introduction

Photoperiod (day length) is a reliable cue by which organisms determine the time of year and coordinate adaptations to seasonal changes in their environment (Goldman, 2001). In nontropical environments, energy-conserving, adaptive adjustments occur among individuals in response to decreasing day lengths that are believed to promote survival during the harsh conditions of winter (Bronson, 1985). For example, short days decrease reproductive system size and function in white-footed mice (Peromyscus leucopus) (Whitaker, 1940; Ruf et al., 1997).

In addition to reproductive involution, individuals of several species also decrease brain mass or specific brain region volume in winter (nonbreeding season) compared with summer (breeding season) (Yaskin, 1984; Smith et al., 1997; Tramontin et al., 1997; Tramontin and Brenowitz, 1998). Reduced hippocampal mass and volume have been reported in short-day rodents (Yaskin, 1984; Perrot-Sinal et al., 1998; Galea and McEwen, 1999). Furthermore, deficits in spatial learning and memory (behaviors that require an intact hippocampus) (Morris et al., 1982) have been reported in short-day relative to long-day rodents (Perrot-Sinal et al., 1998). Spatial memory performance may be decreased during the nonbreeding season when home range size is smaller and roaming to find mates is decreased compared with during the breeding season (Stickel, 1968).

We hypothesized that photoperiod triggers morphological changes in the brain that underlie seasonal adjustments in behavior to promote energetic savings. This hypothesis is supported by spatial learning and memory research in food-storing and brood-parasitic birds and other species (Sherry et al., 1992; Healy et al., 2005).

The hippocampus is among the most plastic sites in adult mammalian brains (McEwen, 1999; Breedlove and Jordan, 1996b). In songbirds, seasonal changes in size and morphology of the brain circuitry mediating song is well established (Brenowitz et al., 1997; Smith et al., 1997; Tramontin et al., 2001). However, studies of seasonal plasticity of mammalian brain structure and function are sparse (for review, see Hofman and Swaab, 2002). Much of the work in rodents has focused on sexual dimorphisms of specific brain regions (Gaulin and FitzGerald, 1989; Jacobs, 1996a).
Changes within the hippocampus include fluctuations in dendritic complexity, spine density, soma size, neurogenesis, and apoptosis and may be associated with altered learning and memory performance (Moser et al., 1994; Woolley, 1998; O’Malley et al., 2000). The goal of this study was to test the hypothesis that short days would decrease hippocampal dendritic complexity and spine density in addition to specifically impairing spatial learning and memory.

We established that short days reduced brain size and hippocampal volume and confirmed previously reported short-day deficits in spatial learning and memory. Additionally, we discovered that photoperiod treatment affected neuronal spine density in the hippocampus. Overall, these data suggest that the functional learning and memory deficits exhibited by short-day mice are associated with structural changes in the hippocampus and reveal that adult mammalian brains display seasonal plasticity.

Materials and Methods

Experiment 1

Animals

Twenty adult male (>55 d of age) white-footed mice (P. leucopus) from a breeding colony maintained at Ohio State University were used in this study. Breeder mice were originally obtained from the Peromyscus Genetic Stock Center at the University of South Carolina (Columbia, SC). All mice were housed in polypropylene cages (27.8 × 7.5 × 13 cm) with a constant temperature and humidity of 21 ± 5°C and 50 ± 10%, respectively, and ad libitum access to food (8640 rodent diet; Harlan Teklad, Indianapolis, IN) and filtered tap water. Mice were housed either in long photoperiods (long-day; n = 7) with a reversed 16:8 h light/dark cycle [lights on at 11:00 P.M. Eastern Standard Time (EST)] or in short photoperiods (short-day; n = 13 males) with an 8/16 h light/dark cycle [lights on at 7:00 A.M. EST] for 13 weeks. Mice were undisturbed except for routine cage changes and biweekly body mass measurements until week 10, when behavioral testing commenced. Mice that failed to regress their gonads in response to short days were dropped from the study (n = 4).

All studies were conducted with approval of the Ohio State Institutional Animal Care and Use Committee approval and in compliance with all United States federal animal welfare requirements.

Sensory tests

Mice were examined for potential photoperiod-induced differences in sensory discrimination before water-maze training. Visual placing was assessed by suspending mice by their tail and noting the first observable extension of forelimbs when lowering mice onto a wire grid from a height of 30 cm (Hatcher et al., 2001). Mice were scored from 0 to 4 for forearm extension (0, none; 1, on nose contact; 2, on vibrissa contact; 3, before vibrissa contact; and 4, early vigorous extension). Tactile discrimination was determined by the presence or absence (scored as 1 or 2, respectively) of a directed head movement in response to brushing a cotton-tipped applicator on the underside of the whiskers. Olfactory and auditory discriminations were tested after the water-maze training at the beginning of the dark phase (3:00–6:00 P.M.). For the olfactory testing, a modified “find the cookie” paradigm was used. Individual mice were placed in a clean cage for 3 min while a 1 cm piece of fruit (orange) was buried in the corner of the home cage. Mice were then returned to their home cages, and latency to uncovering the fruit was recorded for up to 300 s. For auditory discrimination, acoustic startle was tested using a single ventilated startle chamber (SR-Lab; San Diego Instruments, San Diego, CA). Mice were placed inside a clear, nonrestrictive Plexiglas cylinder within the chamber that was hooked up to an electrostatic sensor that measured the amplitude (in millivolts) of displacement caused by the animal’s movement. The mice were subjected to a series of acoustic startle trials (120 dB pulse for a duration of 40 ms) with an average of 15 s between each trial. The average amplitude (in millivolts) transduced by the animal’s movement was used to determine the acoustic startle response. Before onset of the testing session, mice were allowed to acclimatize to the apparatus for 10 min.

Learning and memory tests

Morris water maze. The water maze was used to test long-term spatial learning and memory (Morris, 1984). Testing occurred during the light phase (between 7:00 and 11:00 A.M.). The maze consisted of a white tank (1.3 m diameter) filled with 27°C water to a depth of 47.5 cm. The water was made opaque with white nontoxic tempera paint. The maze was divided into four equal quadrants, and release points were designated at each quadrant as north, east, south, and west. Fixed extramaze cues in the form of large black geometric shapes surrounded the tank. A tracking video camera was suspended from the ceiling above the pool, and 2020 PLUS tracking software (HVS Image, Buckingham, UK) was used. Mice were handled using a small fishing net to avoid the stress of direct handling. On day 1, mice were allowed to swim freely for 60 s without a platform to acclimate to the pool. On days 2–9, a platform (9 cm diameter) was hidden 0.5 cm below the water surface in one quadrant. Mice were given 60 s to locate the hidden platform from random release points around the pool to “escape” from the water three times per day. On reaching the platform or after 60 s, mice were placed on the platform for 10 s and then returned to the home cage. The intertrial intervals were 15 s during which the pool was skimmed of debris. Latency to reach the platform, the distance of the mouse’s path, and swim speed were recorded by the system for each trial to assess acquisition of the spatial task. After the third trial of the day, each mouse received a piece of tissue paper in their cage to expedite drying. On day 10, the platform was removed, and a second probe trial was examine retention. The percentage of time spent in each quadrant (including the quadrant in which the platform had been) was recorded. To evaluate reversal learning, the platform was repositioned in a different quadrant, and retraining to the new location on days 11–15 was completed as described previously. A second probe trial followed reversal training on day 16. On day 17, a single 60 s visible-platform trial was run to determine general visual acuity of the mice in this paradigm. The visual platform (9 cm diameter) was raised 0.5 cm above water level and encircled with a black rim. Latency to reach the platform was recorded.

Spontaneous alternation in a Y maze. On completion of the water-maze task, spontaneous alternation behavior was recorded to assess working memory performance with a spatial component during the dark phase (Crawley, 2000). The maze was made of black Plexiglas and covered with clear Plexiglas to prevent escape during testing. Each arm was 35 cm and converged at an equal angle. Mice were placed in one arm and allowed to move freely through the maze during a 3 min session. The series of arm entries was recorded. An alternation was defined as entries into all three arms on consecutive occasions. The percentage of alternation was calculated using the following formula: (actual alternations/maximum possible alternations) × 100. The maximum possible number of alternations was calculated as the total number of arm entries – 2.

Passive avoidance. Twenty-four hours after spontaneous alternation testing, mice were tested for cued and contextual conditioning memory (nonspatial) using passive avoidance during the dark phase. On day 1 of testing, mice were placed in one chamber of a dark two-chambered box separated by a door (San Diego Instruments). After 30 s of acclimation, a house light in the chamber in which the mouse resided was turned on, and the door was opened. To escape the light, mice entered the other chamber (which remained dark) where they received a 0.06 mA electrical shock (1 s duration) from the grid floor. Mice were then returned to their home cage. On day 2, the same procedure was followed, except the shock was removed. In general, mice that learn to avoid the location of the aversive shock are slow to enter the dark chamber. Latency to enter the dark chamber was recorded to a maximum of 300 s.

Tissue collection and histology

Twenty-four hours after the completion of all behavioral testing, mice were rapidly decapitated. Paired testes were removed and weighed to determine reproductive responsiveness to photoperiod treatment. The average testes mass for long-day mice was determined, and short-day males with a testes mass 2 SD below this mean were considered reproductively responsive to short days and excluded from the study (n = 4). Brains were removed and processed for Golgi staining according to the manufacturer’s protocol (FD Rapid GolgiStain kit; FD Neurotechnolo-
gies, Ellicott City, MD). Throughout the hippocampus, brains were cut coronally in 80 μm sections on a cryostat and mounted on 3% gelatin-coated slides before staining. Sections were also counterstained with cresyl violet. Granule cells in the dentate gyrus (n = 10 per mouse) and pyramidal cells in the CA1 and CA3 regions (n = 5–10 per mouse) were traced using a camera lucida at 400× magnification (NeuroLucida; MicroBrightField, Williston, VT). Neurons chosen for tracing met the following criteria: (1) they were completely impregnated with Golgi stain, (2) they were not obscured by other impregnated neurons or precipitant, and (3) all dendrites were visible within the plane of focus. The hemispheric location of the traced neurons was chosen randomly. Additionally, dendritic spines were traced on five 10 μm distal dendritic segments of each neuron at 1000× magnification of randomly chosen granule cells (throughout the stratum molecular) and pyramidal cells [throughout the stratum oriens (basilar) and the stratum lacunosum-moleculare (apical) that had at least one branch point. Using the accompanying software (NeuroExplorer: MicroBrightField), dendritic complexity (via Sholl analysis), dendritic length, cell body area, and spine density were calculated. The Sholl analysis defines dendritic complexity by the number of dendritic branch points at fixed intervals from the cell bodies (Sholl, 1956). All samples were number-coded, and the experimenter was blind to the treatments.

**Experiment 2**

Sixteen adult male white-footed mice were housed in either long days (n = 8) or short days (n = 8) for 14 weeks. Mice were undisturbed except for routine cage changes and biweekly body mass measurement. At week 14, mice were deeply anesthetized with sodium pentobarbital (Nembutal; Abbott Laboratories, North Chicago, IL) and transectially perfused with 50 ml of saline followed by 75 ml of 10% phosphate-buffered formalin. Brains were removed and postfixed in formalin for at least 24 h at room temperature and then transferred to 15% sucrose in formalin at 4°C overnight. Finally, brains were transferred to 30% sucrose in 0.1 M PBs at 4°C until permeated. Brains were frozen on dry ice and punctured with three stickpins on a rostrocaudal plane through the medial ventral area of the brain. These pinholes were used as reference points to orient the traced sections for volume reconstruction. Brains were then stored at −70°C until being cut in 35 μm sections on a cryostat at −22°C. Every sixth section throughout the hippocampus (from bregma −1.06 to −2.92) (Franklin and Paxinos, 1997) was dry-mounted on 3% gelatin–water (long-day, 539.8 ± 6.5 mg; short-day, 509.6 ± 16.0 mg (p < 0.05 in all cases). Photoperiod treatment had no effect on visual, tactile, olfactory, or auditory discrimination (Table 1) (p > 0.05 in all cases).

**Short days impair Morris water-maze performance**

Short days increased the latency to reach the hidden platform and the distance traveled to reach the hidden platform after seven and eight blocks of trials in the water-maze test (Fig. 1A,B) (p < 0.05 in all cases). The overall effect of photoperiod on latency and path length taken to reach the hidden platform were statistically non-significant (p = 0.08 and 0.1, respectively). However, all animals displayed some degree of learning because both the latency and distance to reach the platform decreased over time (p < 0.001 in both cases). Swim speed, a possible indication of motivation, did not differ at any time point in the hidden-platform trials (data not shown). All mice spent more time in the quadrant in which the platform had been removed for the probe trial (data not shown) (p < 0.05); however, there were no differences between long-day- or short-day-treated mice (p > 0.05).

![Figure 1](image_url) **Figure 1.** Photoperiod-induced differences in spatial learning (via water maze). A, Latency to reach the hidden platform by block of training. B, Path length to reach the hidden platform by block of training. Each block represents three trials. Long-day, n = 7; short-day, n = 9. *p < 0.05.

**Results**

**Short days decrease tissue masses**

Exposure to short days for 13 weeks decreased body mass (long-day, 21.3 ± 0.8 g; short-day, 19.5 ± 0.7 g), testes mass (long-day, 203.5 ± 11.0 mg; short-day, 74.1 ± 9.0 mg), and brain mass (long-day, 539.8 ± 6.5 mg; short-day, 509.6 ± 16.0 mg (p < 0.05 in all cases). Photoperiod treatment had no effect on visual, tactile, olfactory, or auditory discrimination (Table 1) (p > 0.05 in all cases).

**Table 1. Mean ± SEM behavioral scores after 10 weeks of long or short days**

<table>
<thead>
<tr>
<th>Sensory test</th>
<th>Long-day</th>
<th>Short-day</th>
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<tbody>
<tr>
<td>Visual discrimination</td>
<td>2.1 ± 0.1</td>
<td>2.4 ± 0.1</td>
</tr>
<tr>
<td>Vibriosa discrimination</td>
<td>1.5 ± 0.2</td>
<td>1.4 ± 0.1</td>
</tr>
<tr>
<td>Olfactory latency (s)</td>
<td>171.3 ± 30.8</td>
<td>172.6 ± 36.4</td>
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<tr>
<td>Auditory discrimination (% startle)</td>
<td>45.1 ± 7.6</td>
<td>33.1 ± 3.5</td>
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Statistical analyses

Repeated-measures ANOVAs were used to compare water-maze performance over time. Within days, pair-wise comparisons were planned a priori in the analysis models and conducted using two-tailed Student’s t-tests (Keppel, 1991). Student’s t-tests were also used for other behavioral and physiological comparisons. Data with unequal variances were compared using nonparametric Mann–Whitney tests to compare photoperiod differences. One-tailed t-tests were used to compare hippocampal volume because of our a priori hypothesis. All comparisons were considered statistically significant when p < 0.05. StatView version 5.0.1 software was used for all analyses (SAS Institute, Cary, NC).
ing, although this difference did not reach statistical significance \((p = 0.1)\).

**Short days do not affect nonspatial learning and memory**

Photoperiod treatments did not affect the percentage of spontaneous alternation or the total number of possible alternations (a measure of locomotor activity) \((data \ not \ shown)\) \((p > 0.05 \ in \ both \ cases)\).

Both long-day and short-day mice hesitated before stepping through to the side of the chamber that previously shocked them in the passive avoidance test \((day \ 1: \ long-day, 12.3 \pm 4.1 \ s; \ short-day, 6.3 \pm 0.9 \ s; \ day \ 2: \ long-day, 43.6 \pm 29.2 \ s; \ short-day, 24.3 \pm 9.9 \ s)\). However, only the short-day mice hesitated to step over significantly longer on day 2 \((p < 0.05)\).

**Short days affect spine density**

Photoperiod treatment did not affect dendritic complexity in the CA1, CA3, or dentate gyrus regions \((Fig. \ 3A–C)\) \((p > 0.05 \ in \ all \ cases)\). Raw dendritic length also did not differ between photoperiod-treated groups in any of the hippocampal regions examined \((data \ not \ shown)\) \((p > 0.05 \ in \ all \ cases)\). However, short days decreased spine density on the terminal tips \((stratum \ lacunosum-moleculare)\) of both CA1 \((p < 0.05)\) and CA3 \((p = 0.1)\) pyramidal cells compared with long days \((Fig. \ 4A,C)\). On the other hand, short days increased spine density on basilar tips \((stratum \ oriens)\) of CA3 dendrites \((Fig. \ 4B)\) \((p < 0.05)\) but had no effect on spine density in basilar dendritic tips \((stratum \ oriens)\) of pyramidal cells in the CA1 or dendritic tips \((stratum \ moleculare)\) in the dentate gyrus \((p > 0.05)\). Cell body area did not differ between photoperiods in the CA1, CA3, or dentate gyrus regions \((data \ not \ shown)\) \((p > 0.05 \ in \ all \ cases)\).

**Short days decrease hippocampal volume**

Short days decreased both absolute \((long-day, 4.50 \pm 10^{-10} \pm 9.4 \pm 9 \ \mu m^3; \ short-day, 2.86 \pm 10^{-10} \pm 3.3 \pm 9 \ \mu m^3)\) and relative \((hippocampus \ volume/telencephalon \ volume; \ long-day, 1.07 \pm 0.7; \ short-day, 0.22 \pm 0.1)\) hippocampal volume compared with long days \((p < 0.05 \ in \ both \ cases)\). There were no differences in the volume of the telencephalic region surrounding the hippocampus between the photoperiod-treated groups \((long-day, 10.2 \pm 10^{-1} \pm 3.4 \pm 10^{-1} \ \mu m^3; \ short-day, 14.1 \pm 10^{-1} \pm 1.9 \pm 10^{-1} \ \mu m^3)\) \((p > 0.05)\).

**Discussion**

In this study, we observed that short days impaired both spatial-specific learning and memory relative to long days without affecting sensory abilities. Specifically, short-day mice required longer to find a hidden platform in the water maze and spent less time in the quadrant with the platform compared with long-day mice. Furthermore, photoperiod treatment induced changes in hippocampal morphology such that short days decreased apical spine density in the CA1 region of the hippocampus and increased basilar spine density in the CA3 region compared with long days.

We hypothesized that, in addition to inhibiting the reproductive system, short days would trigger morphological changes in the hippocampus that mediate seasonal deficits in learning and memory. The short-day impairment of spatial learning and memory in the water maze supported previous findings in non-breeding deer mice \((Peromyscus \ maniculatus)\) \((Galea \ et \ al., \ 1994; \ Perrot-Sinal \ et \ al., \ 1998)\). We also showed that reversal learning and memory for the reversal platform location are compromised.
Selective impairment of spatial memory in short days
We also tested whether spatial memory is specifically impaired compared with other types of memory based on the hypothesis that spatial memory deficits are derived from the shrinkage of the P. leucopus home range during the nonbreeding season (Stickel, 1968). The selective impairment of long-term spatial learning and memory is supported based on our results from the spontaneous alternation and passive avoidance tests. Additional tests of this hypothesis require experiments using multiple long-term spatial tasks (e.g., radial arm maze). However, controls for photoperiod-induced differences in motivation to eat and locomotion are difficult to equalize in these tests that are dependent on food motivation.

Photoperiodic differences in spine density
In rats, spine density of the dentate gyrus and CA1 region of the hippocampus has been positively correlated with water-maze learning and memory performance (Moser et al., 1994; O’Malley et al., 2000). Spines increase the surface area of dendrites, which increase the excitatory synaptic density and number of connections between neurons (for review, see Sorra and Harris, 2000). Therefore, hippocampal spine density may be an indication of the efficiency of the synaptic network involved in spatial learning and memory. Similarly, our results indicate that apical spine density decreased in the CA1 pyramidal cells of short-day mice in conjunction with their impaired behavioral performance. In contrast to the impaired water-maze performance, however, basilar spine density increased in the CA3 region of short-day mice. The role of spine density among hippocampal regions and dendritic poles in learning and memory remains unspecified (Sorra and Harris, 2000). Photoperiod-induced changes in specific spine morphology remain to be characterized and may provide insight into the functional significance of the present alterations of spine density.

In addition to photoperiod, spine density in adult rodents is also affected by other environmental factors. For example, environmental enrichment (Diamond et al., 1976; Rampon et al., 2000) and awakening after hibernation (Popov et al., 1992) increase spine density in the hippocampus, whereas restraint stress decreases hippocampal spine density (Magarinos et al., 1999). These environment-induced changes in hippocampal spine density occur over varying amounts of time (2 h to 25 d). The time course over which the present photoperiod-induced changes in spine density are established needs to be examined. Neurogenesis in the dentate gyrus, another example of adult hippocampal plasticity, may also be affected by photoperiod (Huang et al., 1998; Lavenex et al., 2000). Studies that test the hypothesis that differences in hippocampal neurogenesis between long-day and short-day mice modulate spatial learning and memory are under way in our laboratory.

Photoperiodic differences in brain and hippocampal size
Gross brain masses were low in voles and shrews trapped during the winter and in juvenile voles exposed to short days from birth (Yaskin, 1984; Dark et al., 1987). In the present study, short days decreased gross brain mass in adult male white-footed mice, both in absolute terms and after correcting for reduced body mass. Additionally, short-day mice had decreased hippocampal volumes, which is consistent with the hypothesis that polygynous (having multiple female mates) rodents increase spatial memory during the breeding season to acquire multiple mates, as well as the hypothesis that hippocampal volume is positively correlated with hippocampal function (i.e., spatial learning and memory) (Gaulin and FitzGerald, 1986; Jacobs et al., 1990; Perrot-Sinal et al., 1998). Similarly, large seasonal variations in home range size have been described in field studies on Peromyscus such that home range sizes are dramatically smaller in the winter (non-breeding season) relative to the summer (Metzgar, 1978). It is important to note that all previously published estimations of hippocampal volume were based on a formula designed to calculate the volume of a truncated cone as opposed to the more sensitive method of volume estimation used in this study (i.e., Cavalieri’s method). In summary, we have linked short day-induced deficits in spatial learning and memory with whole brain and hippocampal involution. These data support the concept that the relative size of a brain region represents its functionality and is capable of displaying adaptive plasticity among seasons in adults (Tramontin and Brenowitz, 2000).

Potential role of hormones on photoperiodic brain changes
Because the duration of melatonin secretion is directly modulated by photoperiod and mediates many photoperiod-induced adaptations, melatonin is a likely hormonal candidate for a mechanism mediating the photoperiod-induced hippocampal modifications described in this report. Melatonin receptors are present in the hippocampal area (entorhinal cortex) of white-footed mice (Weaver et al., 1990; Musshoff et al., 2002). The only model of seasonal plasticity in which the role of melatonin has been examined is in the neural song circuitry of songbirds (Bentley et al., 1999). However, melatonin also affects long-term potentiation in hippocampal slices of photoperiod nonresponsive animals (El-Sherif et al., 2003). Future study of the role of melatonin in seasonal hippocampal plasticity is warranted.

Decreased circulating testosterone and corticosterone concentrations in short-day white-footed mice (Feist et al., 1988) may also modulate morphological hippocampal alterations and result in the learning and memory behavior differences observed in this study. Although breeding condition has been associated with spatial learning and memory performance, the relationship between adult testosterone concentrations and spatial performance is not clear (Galea et al., 1995; Perrot-Sinal et al., 1998), although there appears to be a positive correlation between testosterone concentration and hippocampal volume (Perrot-Sinal et al., 1998; Galea et al., 1999). Additionally, sustained (i.e., chronic) corticosterone treatment or stress-induced increases in corticosterone inhibit learning and memory (for review, see Roozendaal, 2000) as well as decrease hippocampal dendritic complexity (for review, see McEwen and Magarinos, 1997) in various species. We are currently testing the possibilities that photoperiodic alteration of testosterone or corticosterone (basal or stress-induced via the water maze) concentrations mediate the changes described in the present study.

Taken together, the influence of the photoperiod on brain and behavioral plasticity in adults is likely profound in animals that reside in environments with dramatic seasonal changes. Before this study, the link between seasonal effects on brain structure and related function was primarily confined to the neural song circuitry in birds. We suggest that a parallel mammalian model of adult brain plasticity exists in the hippocampus and affects spatial learning and memory. Specifically, morphological changes in the hippocampus appear to underlie seasonal alterations in learning and memory performance. Future research investigating the mechanisms underlying this seasonal plasticity may be applicable to treatment of seasonal cognitive and affective disorders.