Steroid hormones such as estrogens and androgens affect the nervous system during in utero development and throughout adolescence, adulthood, and aging (Choleris et al., 2018). Steroid hormones have been associated with a prevalence in either men or women of certain neurologic disorders. For instance, Alzheimer’s disease and schizophrenia affect more women than men and vice versa, respectively (Irvine et al., 2012; Jackson et al., 2013). Studies have also shown that 17β-estradiol (E2), the predominant and most potent form of estrogen, can alter hippocampal neuroplasticity and therefore affect learning and memory. Exogenous E2 administration into the dorsal hippocampus (DH) of young female rodents enhances memory consolidation as shown by object recognition (OR) and object placement (OP; also known as object location) tasks (Tuscher et al., 2015). OR and OP are two object-based one-trial learning tasks that capitalize on rodents’ innate interest in novelty and relate to the “what” and “where” aspects of episodic memory consolidation and spatial memory, respectively (Frick et al., 2018a). DH E2 infusion immediately after the OR/OP training period restricts the effects of E2 to the time frame of memory consolidation (Tuscher et al., 2015). Post-training DH E2 infusion leads to an increase in spine density of CA1 pyramidal neurons during the time frame in which memory consolidation occurs (Gould et al., 1990; Woolley et al., 1990; Woolley and McEwen, 1993; MacLusky et al., 2005; Phan et al., 2012). Increased dendritic spine density in response to post-training E2 infusion was also observed in the medial prefrontal cortex (mPFC; Tuscher et al., 2016), a structure that expresses estrogen receptors and is also involved in memory consolidation. This observation prompted Tuscher and colleagues to examine in their eNeuro publication the effects of local infusion of E2 into the mPFC and the requirement for concurrent activity of the mPFC in E2-mediated memory consolidation in the DH.

Young female mice were ovariectomized (OVX) to eliminate the ovarian source of E2 that fluctuates during the estrous cycle, the rodent version of the menstrual cycle (Tuscher et al., 2015). Figure 1 shows a schematic illustration of the experimental approach. For OR training, a mouse was placed in a white square arena in which the upper left and right corners contained identical objects, one in each corner (Fig. 1A). The time spent exploring each object was recorded using tracking software, and mice remained in the arena until they had accumulated 30 s exploring the objects. If the mice remembered the identity of the training objects during testing 48 h later, then they should spend more time than chance (>15 s) exploring a novel object that had replaced one of the original training objects in the upper corners of the arena (Fig. 1A). The OP assessment differed from that of OR in two aspects: (1) OP testing occurred 24 h after training; and (2) whereas one of the training objects encountered during training remained in its original location during testing, the second training object had been moved to a lower corner of the arena (Fig. 1B). Spending more time than chance with the moved object indicated that spatial memory was intact.

OVX mice were implanted with bilateral cannulae targeted to the mPFC one week before training in the OR and OP tasks, which were performed in separate groups of mice. Immediately after training, mice received mPFC infusions of either cyclodextrin-encapsulated E2 dissolved in sterile saline or the vehicle control, 2-hydroxypropyl-β-cyclodextrin (HBC) dissolved to the same concentration.
of cyclodextrin as the E2-containing solution. In both the OR and OP behavioral assessments, E2-infused mice spent more time exploring the novel or moved objects than would be expected by chance alone and significantly more time compared to vehicle-infused mice (Fig. 2). These observations showed that local effects of E2 within the mPFC replicate the enhancement of memory consolidation and spatial memory that resulted from local infusion of E2 within the DH.

Although a causal link between E2-mediated spinogenesis and memory consolidation has not yet been established (Sheppard et al., 2019), signaling mechanisms are common to both processes (Frick et al., 2018a). Therefore, Tuscher and colleagues examined local E2-mediated effects on spinogenesis in the mPFC. Two weeks after the completion of the OR and OP behavioral assessments, mice received an infusion of E2 or vehicle control into the mPFC; 2 h later the brains of the mice were harvested and processed for Golgi staining. Dendritic spine analysis was performed in regions in which increased E2-mediated spinogenesis was previously observed, namely, Layer II/III of the mPFC and CA1 of the DH. Spine density was increased apically but not basically when E2 was locally infused into the mPFC, but spine density in the DH was unchanged. It is not known why local infusion of E2 did not affect spinogenesis in the DH; perhaps the time frame was insufficient to detect any alterations or E2 infusion directly into the DH has more hierarchical, rather than reciprocal, effects on another brain region.

The observations described thus far provided evidence that E2 acting locally in the mPFC enhances memory consolidation and increases local spinogenesis. Next, the authors sought to determine if concurrent activation of the mPFC is required for enhanced E2-mediated, DH-dependent memory consolidation. To do so, they employed a designer receptors exclusively activated by designer drugs (DREADD) approach to suppress activity in the mPFC while simultaneously infusing E2 into the DH. Immediately after ovariectomy, viral DREADD constructs or saline (sham condition) were infused bilaterally into the mPFC, and cannulae were also placed bilaterally in the DH for future infusions. The AAV-CaMKIIα-eGFP construct induced expression of green fluorescent protein (GFP) in excitatory neurons in the mPFC three weeks later, at the time when behavioral assessments were initiated. At the time of the surgical procedure, another group of OVX mice received bilateral mPFC injections of an AAV-CaMKIIα-HA-hM4Di-IRES-mCitrine viral construct. Binding of the
clozapine-N-oxide (CNO) ligand to the hM4Di DREADD suppresses the activity of the excitatory CaMKIIα-expressing cells, thereby inhibiting the mPFC. After training in the OR or OP tasks, mice received DH infusion of E2 or vehicle control at the same time that CNO was administered systemically to all experimental groups, at a concentration of CNO that does not impair memory consolidation on its own (Tuscher et al., 2018). In both the OR and OP tasks, memory consolidation was enhanced in mice in which E2 was infused directly into the DH but in which mPFC activity was not suppressed (sham + E2, GFP + E2 groups; Fig. 3) compared to experimental groups that received vehicle control (sham + HBC, GFP + HBC, hM4Di + HBC; Fig. 3). Suppression of excitatory signaling in the mPFC blocked the E2-mediated DH-dependent enhancement of memory consolidation (hM4Di + HBC; Fig. 3) showing, for the first time, that mPFC activity is required for memory consolidation when E2 is locally infused into the DH.

The eNeuro publication by Tuscher and colleagues highlights the complexity of E2-mediated memory consolidation processes and raises several new avenues of research. The finding that local E2 infusion into the mPFC affects a cognitive process that is heavily associated with the hippocampus underscores the importance of research into other brain structures involved in cognition that express steroid receptors. The mPFC and DH interaction could be regulated by direct or indirect neural circuitry. It will also be interesting to determine if local effects of E2 within the mPFC involve classic, genomic signaling via estrogen receptors ERα and ERβ or the rapid signaling that results from extracellular ligand binding to the G protein-coupled receptor GPER. Coupling the use of estrogen agonists and antagonists in the DH with chemogenetic suppression of other brain regions will help resolve the interactions between signaling mechanisms and neural circuitry. Such receptor manipulation could also reveal the relevance of local estradiol production within the hippocampus that is unaffected by ovariectomy.

Tuscher and colleagues’ findings in this eNeuro publication may also help resolve discrepancies in the literature regarding the effects of E2 on cognition by emphasizing the need to examine other brain regions involved in memory consolidation. But more complexities exist, such as the effects of steroid signaling in young versus older or female versus male rodent models (Frick et al., 2018b). It was expected that hormone replacement therapy in postmenopausal women would have a protective effect against cognitive dysfunction. However, some of these studies used conjugated equine estrogens, rather than E2, which may explain the lack of beneficial effects (Luine, 2014). Moreover, it has recently been shown in a mouse model of Alzheimer’s disease that E2 fluctuations during the ovarian cycle have adverse effects on cognition (Broestl et al., 2018). Therefore, a plethora of age, sex, and brain region considerations may be needed to fully elucidate new targets for drug development that address steroid-related effects on cognitive functions, such as memory consolidation.

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