Early exposure to steroid sex hormones can masculinize/defeminize neural tissue. For example, in male rodents, the testes elevate plasma testosterone levels both in utero and in neonates. Testosterone can directly act on androgen receptors or can be metabolized to estradiol, activating estrogen receptors (ER). Stimulation of androgen and ER during development can masculinize neural substrates during a brief critical period; this is referred to as the organizational/aromatization hypothesis (1–3). This process, along with neurosteroids and genetic and environmental influences (4), creates sexually dimorphic neural tissue.

Neuron anatomy and physiology can be sexually dimorphic. This includes adult responsiveness to steroid sex hormones such as estradiol. Thus far, studies of development-induced sex differences in estradiol signaling have largely focused on the classical estradiol action of binding to nuclear-localized ER to directly affect gene expression. Although it is established that early hormone exposure can alter future responsiveness to classical estradiol action, it is unclear whether early hormone exposure can also impact future neuron responsiveness to nonclassical estradiol actions, due to membrane-associated ER (5, 6).

Abbreviations: APDC, (2R,4R)-4-Aminopyrrolidine-2,4-dicarboxylate; CREB, cAMP response element binding protein; DHPG, (S)-3,5-dihydroxyphenylglycine; ER, estrogen receptors; MAP2, microtubule-associated protein 2; mGluR, metabotropic glutamate receptor; pCREB, phosphorylated CREB.
Here we test this hypothesis, using a well-studied, sexually dimorphic, nonclassical estradiol action in hippocampal neurons. In female but not male hippocampal pyramidal neurons, estradiol rapidly modulates cAMP response element binding protein (CREB) phosphorylation (7, 8). This occurs via direct coupling of membrane-associated ERα and -β to metabotropic glutamate receptors (mGluR), leading to estradiol-induced mGluR signaling (schematized in Ref. 6). Female rat neonates were injected once daily for 2 d with vehicle, testosterone, the nonaromatizable androgen dihydrotestosterone, or estradiol. Twenty-four hours after the second injection, hippocampal neuron cultures were generated. Using these cultures, we assessed whether hippocampal pyramidal neurons exhibited membrane ER/mGluR signaling to CREB. Neurons from female pups exposed to vehicle exhibited rapid estradiol action, whereas neurons from male pups exposed to vehicle did not. Neurons from female pups neonatally exposed to testosterone lacked rapid estradiol action. In contrast, neurons from female pups neonatally exposed to dihydrotestosterone exhibited rapid estradiol action. Neurons from female pups neonatally exposed to estradiol, however, lacked rapid estradiol action. Direct mGluR signaling to CREB was unaffected by neonatal hormone exposure. Collectively, these findings indicate that aromatization of testosterone to estradiol leads to a masculinization/defeminization process, whereby hippocampal neurons lose membrane-associated ER signaling. Broadly, these results demonstrate that early hormone exposure contributes to sex differences not only in nuclear ER but membrane ER signaling as well.

### Materials and Methods

#### Animals

All protocols were approved by the Animal Care and Use Committee at the University of Minnesota. Female and male Sprague-Dawley rats were born from timed-pregnant females purchased from Harlan (Indianapolis, IN). Animals were housed with their littersmates and dam. On postnatal d 0 and 1, following a well-established protocol (9), female pups received a sc injection (0.1 ml) of either cottonseed oil (vehicle) or oil containing 100 μg testosterone, estradiol, or dihydrotestosterone. Male pups received cottonseed oil. Each group contained two to four pups, and each experiment was replicated at least three times across different dams and litters.

#### Cell culture

On postnatal d 2, animals were killed, and hippocampal neurons were cultured using previously described techniques (8). Cultures were prepared in parallel from male and female pups, or oil- and hormone-injected pups, obtained from the same litter.

### Drugs

Tetrodotoxin, b(-)-2-amino-5-phosphonopentanoic acid, (S)-3,5-dihydroxyphenylglycine (DHPG), 17β-estradiol, and (2R,4R)-4-aminopyrrolidine-2,4-dicarboxylate (APDC) were from Tocris (Ellisville, MO). Testosterone and dihydrotestosterone were from Steraloids (Newport, RI).

### Immunocytochemistry

Immunocytochemistry protocols followed those described previously (8, 10). Neurons (8–9 d in vitro) were incubated in a Tyrode’s solution containing tetrodotoxin (1 μM) and b(-)-2-amino-5-phosphonopentanoic acid (25 μM) at room temperature for 1.5–2.0 h. Cell stimulations were performed as follows: vehicle for 10 min, 1 nM estradiol for 5 min, 50 μM DHPG for 5 min, 10 μM APDC for 5 min, 20 mM K+ for 3 min, estradiol or APDC for 5 min, and then estradiol or APDC and 20 mM K+ for 3 min. Cells were fixed using ice-cold 4% paraformaldehyde (Electron Microscopy Sciences, Fort Washington, PA) in PBS containing 4 mM EGTA. Cells were washed in PBS, permeabilized with 0.1% Triton X-100 (VWR Scientific, West Chester, PA), washed, and blocked at 37°C for 30 min in PBS containing 1% BSA and 2% goat serum (Jackson ImmunoResearch, West Grove, PA). The cells were incubated at 37°C for 1 h in block containing a monoclonal antibody directed against serine 133 phosphorylated CREB (pCREB) (1:1000, 05-667; Upstate Biotechnology, Lake Placid, NY) and a polyclonal antibody targeting microtubule-associated protein 2 (MAP2) (1:1000, AB5622; Calbiochem, La Jolla, CA). Cells were washed and incubated for 1 h at 37°C in block solution containing fluorescein isothiocyanate- and cyanine 5 (Jackson ImmunoResearch) or Alexa Fluor 488-conjugated antirabbit and 635-conjugated antimouse (Invitrogen, Carlsbad, CA) secondary antibodies for visualization of MAP2 and pCREB, respectively. Cells were washed and mounted using Citifluor (Ted Pella, Redding, CA). Nuclear fluorescent intensities for pCREB were acquired using a Leica DM5500Q confocal system or a Yokogawa spinning-disc confocal system. Data acquired from the Yokogawa system were quantified using MetaMorph (version 6.0; Universal Imaging, Downingtown, PA). Data from the Leica were quantified with Leica LAS AF (version 1.9.0; Leica, Buffalo Grove, IL). The same antibodies and imaging system were used for all cells within an experiment. Experimental conclusions were not altered by these differences.

Following established protocols (8), the confocal excitation and detection settings for each experiment were determined using 20 mM K+ -stimulated coverslips. Inter-coverslip variability was accounted for by subjecting two coverslips to each treatment. Neurons were selected randomly across both coverslips using MAP2 fluorescence, allowing blind acquisition of pCREB intensities. Data acquisition order was random. Images were captured through the approximate midline of each cell. During data analysis, MAP2 staining was used to draw a region of interest outlining the nucleus of each neuron, blinding analysis of pCREB intensity. The region of interest was then transferred to the pCREB image, and average fluorescence intensities within the nucleus were recorded. Background from a region of the image not containing pCREB fluorescence was subtracted from the average pCREB fluorescence.
Experiments were analyzed using one-way ANOVAs and Tukey’s post hoc test (Prism version 5.00; GraphPad Software, La Jolla, CA). Statistical differences between groups are depicted within each figure as different alphabetical characters. *P* values < 0.05 were considered *a priori* as significant. Data are presented as mean ± SEM.

**Results**

**Only hippocampal neurons from females exhibit rapid estradiol regulation of CREB phosphorylation**

Estradiol rapidly phosphorylates CREB in female hippocampal pyramidal neurons, and preexposure to estradiol attenuates CREB phosphorylation induced by the depolarizing action of 20 mM K⁺ (8). These rapid estradiol effects are mediated through membrane-associated ER coupling to mGluR and are not observed in neurons from males (7, 8). For our first experiment, we verified that neonatal oil injections would not affect this outcome. We injected male and female littermate rats with vehicle on postnatal d 0 and 1 and then cultured hippocampal pyramidal neurons on d 2. After 8–9 d in culture, we assessed neuronal responsiveness to estradiol, examining changes in CREB phosphorylation using the following treatments: vehicle, estradiol, 20 mM K⁺, or estradiol with 20 mM K⁺ (Fig. 1A). In neurons from female animals [Fig. 1B; *F*(3,116) = 70.38, *P* < 0.0001], exposure to estradiol increased CREB phosphorylation compared with vehicle (*P* < 0.01). Estradiol also attenuated 20 mM K⁺-induced CREB phosphorylation compared with 20 mM K⁺ alone (*P* < 0.01). Neurons from male animals did not respond to estradiol either under baseline or 20 mM K⁺ conditions [Fig. 1C; *F*(3,118) = 47.12, *P* < 0.0001].

**Neonatal exposure to testosterone eliminates rapid estradiol action**

This sexually dimorphic sensitivity to estradiol could be organized early in life by hormone exposure, by a genetic program independent of early hormone exposure, or by environmental factors. To differentiate between these options, we followed the same experimental timeline as above, this time with neonatal injections of vehicle or testosterone. We reasoned that if in males neonatal exposure to masculinizing doses of testosterone eliminates later responsiveness to estradiol, then a similar treatment in females should also eliminate rapid estradiol action.

Neonatal testosterone exposure eliminated rapid estradiol action [Fig. 1D; *F*(3,117) = 155.0, *P* < 0.0001]. In neurons from female animals [Fig. 1B; *F*(3,116) = 70.38, *P* < 0.0001], exposure to estradiol increased CREB phosphorylation compared with vehicle (*P* < 0.01). Estradiol also attenuated 20 mM K⁺-induced CREB phosphorylation compared with 20 mM K⁺ alone (*P* < 0.01). Neurons from male animals did not respond to estradiol either under baseline or 20 mM K⁺ conditions [Fig. 1C; *F*(3,118) = 47.12, *P* < 0.0001]. These results support the hypothesis that membrane-associated ER signaling is lost upon early hormone exposure.
Neonatal exposure to dihydrotestosterone does not eliminate rapid estradiol action

In the brain, testosterone can act directly on androgen receptors or be metabolized into other hormones. These include dihydrotestosterone and estradiol. Dihydrotestosterone is a potent, nonaromatizable androgen that binds to androgen receptors, whereas estradiol binds to ER. To determine whether the effects of testosterone were due to activation of androgen receptors, we again followed the same experimental timeline, with neonatal injections of vehicle or dihydrotestosterone.

Neonatal exposure to dihydrotestosterone did not eliminate rapid estradiol action [Fig. 1E; F(3,118) = 72.97, P < 0.0001]. In neurons from females neonatally exposed to dihydrotestosterone, exposure to estradiol increased CREB phosphorylation compared with vehicle (P < 0.001) and attenuated 20 mM K⁺-induced CREB phosphorylation compared with 20 mM K⁺ alone (P < 0.001)

Similar effects of estradiol were observed in neurons from vehicle-treated animals [F(3,116) = 70.38, P < 0.0001]. These results indicate that the organizational effect of early exposure to testosterone on hippocampal neuron estradiol sensitivity is not mediated via androgen receptors.

Neonatal exposure to dihydrotestosterone does not eliminate rapid estradiol action

Given that testosterone can also be metabolized into estradiol, we next tested the role of early ER activation. We followed the same experimental timeline as above, with neonatal injections of vehicle or estradiol.

Neonatal estradiol exposure eliminated rapid estradiol action [Fig. 1F; F(3,128) = 57.87, P < 0.0001]. In neurons from females neonatally exposed to estradiol, exposure to estradiol did not increase CREB phosphorylation compared with vehicle (P = 0.05) and estradiol did not attenuate 20 mM K⁺-induced CREB phosphorylation compared with 20 mM K⁺ alone (P > 0.05). This is in contrast to the neurons obtained from females exposed to vehicle, which showed normal responses to estradiol [F(3,117) = 25.21, P < 0.0001]. These results support the hypothesis that sex differences in estradiol responsiveness are generated by activation of ER.

mGluR signaling is unaffected by neonatal exposure to testosterone

In both males and females, mGluR activation regulates CREB phosphorylation. Only in females, however, are membrane-associated ER functionally coupled to mGluR (7, 8). The hormone manipulations employed here eliminate this membrane-associated ER signaling in females. Our interpretation of these data is that the early hormone exposure is masculinizing/defeminizing the females by inducing a male-like phenotype: i.e. eliminating membrane-associated ER coupling to mGluR. If this interpretation is correct, then the direct mGluR pathway to CREB would remain intact in the masculinized/defeminized females. To test this hypothesis, we neonatally injected females with either vehicle or testosterone and then assessed neuronal sensitivity to the group I mGluR agonist DHPG and the group II mGluR agonist APDC on d 8 or 9. Neonatal exposure to vehicle did not eliminate ER or group I mGluR signaling [Fig. 2A; F(2,109) = 9.143, P = 0.0002]. In contrast, neonatal exposure to testosterone eliminated estradiol signaling without affecting group I mGluR signaling [Fig. 2B; F(2,108) = 17.34, P < 0.0001]. Similar results were obtained regarding group II mGluR signaling. Neonatal exposure to vehicle did not eliminate estradiol or group II mGluR signaling [Fig. 2C; F(4,146) = 35.80, P < 0.0001]. However, neonatal exposure to testosterone

FIG. 2. Neonatal exposure to testosterone (T) does not affect mGluR signaling. A, Neonatal exposure to oil does not affect group I mGluR signaling induced by the agonist DHPG or rapid estradiol (E) action in female neurons; B, neonatal exposure to testosterone does not eliminate group I mGluR signaling but does eliminate rapid estradiol action (E) in female neurons; C, neonatal exposure to oil does not eliminate group II mGluR signaling induced by the agonist APDC (AP) in female neurons; D, neonatal exposure to testosterone does not affect group II mGluR signaling but does eliminate rapid estradiol action on pCREB intensity induced by 20 mM K⁺. Letters within each bar indicate statistically significantly different groups; complete statistical information is in Results.
eliminated estradiol signaling without affecting group II mGluR signaling [Fig. 2D; $F_{(4,160)} = 47.77$, $P < 0.0001$]. Overall, these data indicate that mGluR signaling to CREB is intact in masculinized/defeminized females, supporting the hypothesis that neonatal hormone exposure specifically modulates membrane ER signaling.

**Discussion**

There are five principle findings of these experiments. First, rapid, nonclassical estradiol modulation of CREB phosphorylation occurs in female but not male hippocampal pyramidal neurons. Second, hippocampal pyramidal neurons from female neonates exposed to testosterone lacked estradiol signaling to CREB. Third, dihydrotestosterone injections of female neonates did not disrupt estradiol regulation of CREB. Fourth, estradiol injections of female neonates eliminated estradiol signaling to CREB. Fifth, masculinization/defeminization does not affect mGluR signaling. Collectively, these experiments demonstrate that the organizational effects of early hormone exposure apply to rapid, nonclassical estradiol action.

Nonclassical estradiol action has been known since at least the 1960s, when Szego and Davis (11) demonstrated that within seconds, estradiol increased cAMP accumulation in uterine tissue. Later, in neurons, Kelly and colleagues (12) showed that estradiol could rapidly affect neuron electrophysiological properties. These discoveries were roughly in parallel with those delineating classical estradiol actions via ER acting in the nucleus to affect gene transcription via estrogen response elements (6, 13). Although initially controversial, today, it is well accepted that estradiol rapidly modulates neuronal electrophysiological properties (14–16). It is also definitive that estradiol can induce second messenger signaling pathways more commonly associated with G protein-coupled receptors to modify cellular physiology, gene transcription, and even anatomy (6, 13, 17–20). Indeed, nonclassical estradiol action can masculinize neural tissue (21). Here, we have chosen to work with rapid, nonclassical estradiol modulation of CREB phosphorylation in hippocampal pyramidal neurons (8). In female but not in male neurons, this modulation occurs via membrane-associated ERα and -β that couple to mGluR (schematized in Ref. 6). This coupling is widespread across the nervous system, including hippocampal (8, 22), striatal (23), cortical (24), arcuate (25), and dorsal root ganglion neurons (26) as well as hypothalamic astrocytes (27).

This association between membrane ER and mGluR, as well as other G proteins, occurs in brain regions that also express nuclear-localized ER that operate via the classical mechanism. This integration allows estradiol to not only slowly change gene expression but also exert influence over the functions typically ascribed to G protein-coupled receptors. This spectrum of estradiol action could thus potentially act in parallel to modulate neuron function (28). Examples of this include the arcuate nucleus/medial preoptic nucleus circuit, where rapid estradiol signaling facilitates classical nuclear ER action via mGluR stimulation (13, 25), experiments using the membrane-ER-knockout mouse, which found that normal development requires both membrane and nuclear ER (29), and teleosts, where rapid steroid hormone-specific modulation of a neural circuit controlling vocal behavior differs between sexes (30).

These complementary actions of classical and nonclassical estradiol signaling suggest that common developmental mechanisms underlie sexual dimorphisms in the diverse spectrum of estradiol action. Supporting this, use of the four core genotypes of mice (31) indicated that the sexually dimorphic nonclassical response of hypothalamic astrocytes to estradiol was dependent upon gonadal but not chromosomal sex (32). This implicates early hormone exposure, foreshadowing the conclusions presented here. It does not necessarily follow, however, that all sexually dimorphic sensitivity to nonclassical hormone signaling is organized by early hormone exposure. This is only one of several possible mechanisms, with others including neurosteroids and genetic and environmental influences (4). Thus, the developmental origin of sexually dimorphic phenomenon must continue to be evaluated on a case-by-case basis.

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