NEONATAL INHIBITION OF BRAIN ESTROGEN SYNTHESIS ALTERS ADULT NEURAL FOS RESPONSES TO MATING AND PHEROMONAL STIMULATION IN THE MALE RAT

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Abstract—Neonatal inhibition of brain estrogen formation in male rats by administration of the aromatase inhibitor, 1,4,6-androstatriene-3,17-dione (ATD), permanently changes aspects of their mating behavior and partner preference in adulthood. The medial preoptic area receives chemosensory inputs via a sexually dimorphic vomeronasal projection circuit, which responds to reproductively relevant pheromonal cues. The medial preoptic area also receives genital somatosensory inputs via the midbrain central tegmental field and the medial amygdala. We used Fos immunoreactivity as a marker of neuronal activation to determine whether there is a correspondence between the behavioral profiles of neonatally ATD-treated male rats and their neuronal responses in the medial preoptic area and other brain regions to somatosensory and chemosensory stimuli.

Achieving eight intromissions with an estrous female led to a greater neuronal Fos immunoreactivity in the medial preoptic area of neonatally ATD-treated male rats compared with neonatally cholesterol-treated male rats. Exposure for 1.5 h to chemosensory cues derived from soiled bedding of estrous females induced Fos immunoreactivity throughout the vomeronasal pathway (i.e. medial amygdala, bed nucleus of the stria terminalis and medial preoptic area) in both ATD and cholesterol males (Experiment 2a). By contrast, exposure for 1.5 h to chemosensory cues derived from soiled bedding of sexually active males revealed clear differences between ATD and cholesterol males in neuronal Fos immunoreactive (Experiment 2b). At peripheral portions of the vomeronasal pathway (i.e. the accessory olfactory bulb and the medial amygdala), there were no differences in the number of Fos-immunoreactive neurons between ATD and cholesterol males. However, neurons in the more central portions of the vomeronasal pathway (i.e. the bed nucleus of the stria terminalis and the medial preoptic area) showed increased Fos immunoreactivity after exposure to odors from sexually active males in ATD males as opposed to cholesterol males. Females, like ATD males, showed neuronal Fos immunoreactivity at each level of the vomeronasal pathway after being exposed to odors from sexually active males.

These results suggest that the responsiveness of neurons in the central portion of the vomeronasal projection circuit to odors from sexually active males, but not estrous females, is sexually differentiated in male rats due to the neonatal action of estrogens. Copyright © 1996 IBRO. Published by Elsevier Science Ltd.

Key words: neonatal ATD treatment, Fos immunoreactivity, sexual behavior, pheromonal stimulation, preoptic area.

Perinatal administration of male rats of 1,4,6-androstatriene-3,17-dione (ATD), which blocks the aromatization of testosterone to estradiol, affects the sexual differentiation of the CNS, as well as psycho-
preference for the sexually active male, whereas normal males preferred the estrous female.3

Numerous studies have shown that bilateral lesions of the medial preoptic area (mPOA) anterior hypothalamus continuum disrupt masculine coital function in male vertebrates and enhance males’ perceptive and receptive responsiveness towards a stimulus male in cats,9 rats,20,24,26 and guinea-pigs,29 and induces a partner preference for a male in ferrets.27 Within the mPOA, a sexually dimorphic nucleus (SDN-POA), which is significantly larger in males than in females, was first described in rats.17 In rats, the sexual differentiation of the SDN-POA shows a close parallel with psychosexual differentiation, i.e. its larger size in males depends on the action of estrogenic metabolites of testosterone during a critical perinatal period.14,15,22 This correlation was emphasized by a recent study in which SDN-POA volume was reduced in both prenatally and pre- and neonatally ATD-treated males, with a larger reduction being achieved in the latter than in the former group.21

Immunocytochemical visualization of Fos, the nuclear protein product of the immediate early gene, c-fos, has provided useful information about the neural circuits which are activated following mating in male rats.5 In male rats, neural Fos immunoreactivity (IR) is increased after mating in several forebrain regions, including the mPOA, bed nucleus of the stria terminalis (BNST) and posterodorsal portion of the medial amygdala (MePD), and in the midbrain central tegmental field (CTF).5,6,28,34 We used Fos-IR as a marker of neural activity to reveal possible differences between neonatally estrogen-deprived (i.e. neonatally ATD-treated) males and normal (i.e. neonatally cholesterol-treated) male rats’ responses to genital and pheromonal stimulation associated with sexual behavior. In Experiment 1, the number of Fos-IR neurons was compared in several brain regions of adult male rats treated neonatally with ATD or cholesterol. Animals were killed 1 h after achieving eight intromissions with an estrous female.

In rodent species, reproductivity relevant pheromonal cues are detected by receptors in the vomeronasal organ, which in turn transmit this information centrally via the accessory olfactory bulb (AOB), the medial amygdala, BNST and mPOA.23 In the rat, this vomeronasal projection pathway is sexually dimorphic, with males having more vomeronasal receptors and a larger number of neurons throughout.3 In male rats, the sexual differentiation of this vomeronasal pathway apparently occurs under the influence of perinatal estrogens.16,31 Therefore, in Experiment 2, we investigated whether pheromonal cues (derived from either the urine and feces of estrous females or sexually active males) may reveal differences in neural c-fos responses in the vomeronasal projection to the mPOA and/or the nucleus accumbens, a brain region which has been linked to reward and sexual motivation,14,25,26 between neonatally ATD-treated males and cholesterol-treated males.

EXPERIMENTAL PROCEDURES

Animals

Male and female Wistar RP rats bred in our laboratory were housed in single-sex groups of two to three. Food and water were available ad libitum. All rats were kept on a reversed 14:10 h light–dark cycle (lights off from 7.45 a.m. to 5.45 p.m.). Female rats were time-mated and parturition occurred 22 days later. Within 2-4 h after birth, the newborn males received subcutaneously a silastic capsule (SR3: inner diameter 1.5 mm, outer diameter 2.1 mm, length 5 mm) containing crystalline ATD or cholesterol under ice anesthesia. The implants were removed at 21 days of age (day of weaning), and the animals were housed two to three of the same treatment to a cage.

Experiment 1: neural c-fos responses to mating

At the age of approximately three months, 15 neonatally ATD-treated and 15 cholesterol-treated males were castrated under ether anesthesia and were injected daily with testosterone propionate (200 µg, s.c., in olive oil). The ATD and cholesterol males were pretreated three times (30 min/test) with an estrous female in order to provide them with heterosexual experience. All behavioral testing was conducted in semicircular cages measuring 62 × 40 × 36 cm. Before testing, the experimental animal spent a 15-min adaptation period in the cage. The test began when a female was put in the cage. The stimulus females were ovarioctomized and brought into behavioral estrus by injecting 20 µg of estradiol benzoate 24–48 h followed by 1.0 mg of progesterone 3–4 h before testing.

All males were housed singly for one or two days prior to the actual experiment. Groups of ATD and cholesterol males were allowed to achieve eight intromissions with an estrous female. Then, these males were put singly in their home cages for 1 h, after which they were perfused and their brains processed for Fos-IR. Pairs of ATD and cholesterol males were matched for number of mounts and intromissions, and testing time. Additional male rats (ATD and cholesterol; housed singly for one or two days) taken directly from their home cages served as unmated controls. In Experiment 1, 10 mated ATD, 10 mated cholesterol, five unmated ATD and five unmated cholesterol males were used.

Experiment 2a: neural c-fos responses to chemosensory cues derived from soiled bedding of estrous females

At the age of approximately six months, five neonatally ATD-treated and 10 cholesterol-treated males were castrated under ether anesthesia using a midline abdominal incision and received estradiol subcutaneously through a silastic capsule (inner diameter 0.5 mm, outer diameter 1.0 mm, length 2.5 cm). We chose adult estradiol treatment because the sexual behavior of ATD males differed most dramatically from that of control males under this hormonal condition.17 Two groups of ovarioctomized females (six per group) were injected subcutaneously with estradiol benzoate (20 µg) 24–48 h prior to a progesterone (1.0 mg) injection. Three hours later, all females were placed in clean cages which contained fresh sawdust. Bedding was collected 12 h later and immediately used in the experiment. The heterosexually experienced ATD and cholesterol males were housed singly for one to two days prior to use in the experiment in a room containing no other rats. Pairs of ATD and cholesterol males were perfused after spending 1.5 h in the cage containing soiled bedding from estrous females and their brains were processed for Fos-IR. Additional male rats, housed singly in a separate room...
Neonatal inhibition of brain estrogen synthesis

Fig. 1. Photomicrographs showing Fos-IR in the mPOA of an unmated cholesterol-treated male (A), a cholesterol-treated male after eight intromissions with an estrous female (B) and an ATD-treated male after eight intromissions with an estrous female (C). All males were castrated in adulthood and subsequently treated with testosterone propionate.

containing no other rats, were taken directly from their home cages to the perfusion room and served as controls not exposed to pheromones (home cage males). Because no differences in c-fos responses were found between the unmated ATD and cholesterol males in Experiment 1, we used only neonatally cholesterol-treated males as home cage males. All experiments took place in the early part of the dark phase of the light–dark cycle (between 8.30 and 11.30 a.m.).

Experiment 2b: neural c-fos responses to chemosensory cues derived from soiled bedding of sexually active males

At the age of approximately nine months, seven neonatally ATD-treated and seven cholesterol-treated males were castrated under ether anesthesia using a midline abdominal incision and received estradiol subcutaneously through a silastic capsule (inner diameter 0.5 mm, outer diameter 1.0 mm, length 2.5 cm). The female rats used in this experiment were ovariectomized and treated with estradiol benzoate 24–48 h prior to the experiment. The ATD and cholesterol males were pretested twice with an estrous female (15 min/test) and twice with an active male (15 min/test) in order to provide them with sexual experience. The females had been used as stimulus animals during partner preference testing earlier in our laboratory.

Three groups of males (six per group) were placed in clean cages which contained fresh sawdust. Bedding was collected 72 h later and immediately used in the experiment. All experimental animals (ATD, cholesterol and females) were housed singly one or two days prior to use in the experiment in a room containing no other rats (females separate from males). The experimental animals were perfused after spending 1.5 h in the cage containing soiled bedding from active males and their brains were processed for Fos-IR. A control group, comprising two ATD males, two cholesterol males and two females, was killed after spending 1.5 h in clean bedding (clean bedding subjects). We chose this particular composition of the control group because: (i) no differences in Fos-IR were found between unmated ATD and cholesterol males in Experiment 1; (ii) neural c-fos responses in

Table 1. Effect of eight intromissions with an estrous female on the mean (±S.E.) number of Fos-immunoreactive neurons (counted per standard area, 0.27 mm²) in neonatally estrogen-deprived (ATD-treated) and normal males (cholesterol-treated)

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<th>Group</th>
<th>n</th>
<th>Core</th>
<th>Shell</th>
<th>mPOA</th>
<th>SDN-POA†</th>
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<tr>
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<td>6±1</td>
<td>20±4</td>
<td>4±2</td>
<td>10±2</td>
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<td>Cholesterol</td>
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<td>52±9*</td>
<td>119±9</td>
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<td>43±5*†</td>
<td>84±8*</td>
<td>104±7*</td>
<td>34±5*</td>
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All animals were castrated and treated with testosterone propionate at the time of the experiment.

*Significantly (P < 0.05) higher compared to unmated males.
†Significantly (P < 0.05) higher in mated ATD males compared to mated cholesterol-treated males.
‡SDN-POA data are expressed as mean number of Fos-IR neurons inside the SDN-POA per mean SDN-POA area.
home cage subjects in Experiment 2a were very low; (iii) females have been added to the experimental groups. All experiments took place in the early part of the dark phase of the light-dark cycle (between 8.00 and 11.00 a.m.).

**Immunocytochemistry**

Rats were anesthetized with sodium pentobarbital (100 mg/rat, i.p.), given an intracardiac injection of heparin (1000 U/rat) and perfused via the aorta with 0.1 M phosphate-buffered saline (PBS; pH 7.3) followed by 4% paraformaldehyde in 0.1 M phosphate buffer (pH 7.3). Brains were removed and postfixed in 4% paraformaldehyde for 2 h before being placed in 20–30% sucrose–PBS for cryoprotection overnight on a shaker at 4°C. Coronal sections of 52 μm were cut using a sledge microtome with a freezing stage. In Experiments 1 and 2a, consecutive sections were saved at each of four levels, including the nucleus accumbens, mPOA/BNST, medial amygdala and midbrain CTF (in Experiment 1 only). In Experiment 2b, consecutive sections were saved at each of three levels, including the olfactory bulb, the nucleus accumbens and from the medial preoptic anterior hypothalamus to the MePD. Free-floating sections were rinsed twice in 0.1 M PBS and then incubated overnight on a shaker at room temperature with an anti-Fos antiserum raised in rabbit against the N-terminal sequence of rat Fos amino acids (DCH-1; a gift from Dr David Hancock, London, U.K.). After incubation in primary antiserum, the brain sections were rinsed four times in 0.1 M PBS containing 0.02% Triton X-100 and incubated for 2 h on a shaker at room temperature with biotinylated goat anti-rabbit immunoglobulin G. After incubation with the secondary antiserum, the sections were rinsed four times and incubated for 1.5 h on a shaker at room temperature with avidin–biotin–peroxidase complex solution (ABC Elite Kit, Vector Laboratories). The brain sections were again rinsed four times in 0.1 M PBS (no Triton X-100) and then reacted with nickel chloride–3,3’-diaminobenzidine and 0.0003% hydrogen peroxide for 5–10 min (DAB Kit, Vector Laboratories). The sections were rinsed three times, mounted onto gelatin-coated slides and coverslipped using Permount. In Experiment 1, after the immunocytochemistry for Fos protein was completed, alternate sections were counterstained with Cresyl Violet in order to determine the boundaries of the SDN.

**Analysis**

To quantify the numbers of Fos-IR nuclei, all slides were coded so that the investigator had no knowledge of sex or treatment condition of brain sections from any subject. In Experiment 1, two brain sections at the level of the nucleus accumbens, mPOA, BNST, MePd and CTF were selected for quantitative analysis for each animal. In Experiment 2a, brain sections at the level of the nucleus accumbens, mPOA, BNST, posteroverentral portion of the medial amygdala (MePV) and MePD were selected for quantitative analysis for each animal. We did not collect the olfactory bulbs in Experiment 2a, because Bressler and Baum reported similar increments in the number of Fos-IR neurons in the olfactory bulb for males and females. Thus, we had no expectation of finding a difference in Experiment 2a. In Experiment 2b, brain sections at the level of the AOB (mitral and granule cell layer), nucleus accumbens, mPOA, BNST, MePV and MePD were selected for quantitative analysis for each animal. All of the Fos-IR nuclei in a field of view under the ×25 objective (0.27 mm²) were drawn using a camera lucida and counted later.

To analyse the distribution of Fos-IR nuclei in the SDN-POA (Experiment 1), the number of Fos-IR nuclei inside the SDN was counted for each animal in the
counter-stained sections. Because the SDN was present in one to three sections only, the mean number of Fos-IR nuclei inside the SDN was calculated for each animal. Area measurements of the SDN (coronal sections) were performed unilaterally using the method described by Houts-muller et al.21 and a mean SDN area was calculated for each animal. The Fos-IR data of the SDN are expressed as mean number of Fos-IR nuclei inside the SDN/mean SDN area. In some animals (two cholesterol unmated, one ATD and one cholesterol mated), the sections taken for the mPOA/BNST did not include the SDN. These animals were excluded from statistical analysis. In addition, camera lucida drawings of Fos-IR nuclei and the SDN were made of consecutive coronal sections through the mPOA of a representative ATD male and a cholesterol male after mating.

**Statistics**

All Fos-IR data were first analysed for normal distribution using the Kolmogorov–Smirnov test and were subsequently subjected to two-way ANOVA in Experiment 1 (neonatal treatment x mating) or one-way ANOVA in Experiment 2 (groups), followed by the Student–Newman–Keuls method for pairwise multiple comparison procedures (Student–Newman–Keuls method). If the normality test failed (P < 0.05), the data were subjected to Kruskal–Wallis one-way ANOVA on ranks, followed by the Student–Newman–Keuls method for pairwise multiple comparison procedures. In Experiment 1, the sections containing the nucleus accumbens were taken too caudal for some animals (two ATD and one cholesterol unmated, one ATD and one cholesterol mated). These animals were excluded from statistical analysis.

**RESULTS**

**Experiment 1: neural c-fos responses to mating**

Mounting and intromitting with an estrous female significantly augmented the number of Fos-IR neurons in several brain regions of both ATD and cholesterol males. The magnitude of this c-fos response was very similar in ATD and cholesterol males, with the exception of the response in the mPOA, which was greater in ATD males than in cholesterol males. An example of the effect of mating on Fos-IR in the mPOA is shown in Fig. 1. Two-way ANOVA showed that there was a significant effect of mating \[ F(1/26) = 80.9, P < 0.0001 \] and a significant effect of neonatal treatment \[ F(1/26) = 5.9, P = 0.02 \] on the number of Fos-IR nuclei in the mPOA (Table 1). Post hoc analyses revealed that there were significantly more Fos-IR nuclei in the mPOA of mated males than of unmated males. Furthermore, mated ATD males had significantly more Fos-IR nuclei in the mPOA than mated cholesterol males. Mating also augmented the number of Fos-IR nuclei in the BNST \[ F(1/26) = 57.2, P < 0.0001 \], MePD \[ F(1/26) = 72.8, P < 0.0001 \] and CTF \[ F(1/26) = 21.9, P < 0.0001 \] (Table 1) to equivalent degrees in ATD and cholesterol males. Finally, the nucleus accumbens shell \[ F(1/20) = 28.2, P < 0.0001 \] and core \[ F(1/20) = 15.0, P = 0.001 \] also showed increased Fos-IR after mating in both ATD and cholesterol males (Table 1).

The mating-induced increments in Fos-IR occurred in portions of the mPOA which included, but were not restricted to, the SDN. Camera lucida
drawings of Fos-IR neurons and the SDN in consecutive coronal sections through the mPOA of mated males are shown in Fig. 2. Two-way ANOVA (neonatal treatment × mating) on the mean number of Fos-IR neurons inside the SDN (see Table 1) showed a significant effect of mating \[F(1/22) = 29.4, P < 0.0001\] and of neonatal treatment \[F(1/22) = 4.2, P = 0.05\] on the number of Fos-IR nuclei in the SDN. Post hoc analyses revealed that the number of Fos-IR nuclei was significantly higher in mated males than in unmated males. Furthermore, the SDN of mated ATD males contained more Fos-IR neurons than the SDN of mated cholesterol males. There were no significant differences between ATD and cholesterol males in mean SDN area (ATD: 0.066 ± 0.005 mm²; cholesterol: 0.078 ± 0.007 mm²).

Experiment 2a: neural c-fos responses to chemosensory cues derived from soiled bedding of estrous females

Exposure to soiled bedding of estrous females significantly augmented the number of Fos-IR nuclei at each level of the vomeronasal projection circuit of both ATD and cholesterol males, compared to home cage control males. The magnitude of this c-fos response was very similar in ATD and cholesterol males. One-way ANOVA revealed that exposure to pheromonal cues of estrous females augmented the number of Fos-IR neurons in the MePV \[F(2/12) = 8.1, P = 0.006\] and MePD \[F(2/12) = 18.6, P = 0.0002\] to equivalent numbers in ATD and cholesterol males (Table 2). Pheromonal stimulation also increased Fos-IR in the BNST \[F(2/12) = 8.1, P = 0.006\] and mPOA \[F(2/12) = 15.1, P = 0.0005\] similarly in ATD and cholesterol males (Table 2). Furthermore, there was a significant pheromonal-induced increment in Fos-IR in the nucleus accumbens core \[F(2/12) = 5.9, P = 0.02\] and shell \[F(2/12) = 4.2, P = 0.04\] to equivalent degrees in ATD and cholesterol males (Table 2).

Experiment 2b: neural c-fos responses to chemosensory cues derived from soiled bedding of sexually active males

There were clear differences between ATD and cholesterol males in neuronal c-fos expression following exposure to soiled bedding of sexually active males. At peripheral stages of the vomeronasal pathway (i.e. the AOB and MePD), ATD and cholesterol males showed similar c-fos responses, but at more central levels of the vomeronasal pathway (i.e. the

Fig. 3. Photomicrographs of the AOB showing FOS-IR neurons in the mitral (m) and granular (gr) layers of a male after spending 1.5 h on clean bedding (A), a cholesterol-treated male after spending 1.5 h on soiled bedding from sexually active males (B), an ATD-treated male after spending 1.5 h on soiled bedding from sexually active males (C) and a female after spending 1.5 h on soiled bedding from sexually active males (D). All animals were gonadectomized in adulthood and subsequently treated with estradiol.
BNST and mPOA), ATD males still showed clear \(c\)-fos responses, whereas cholesterol males did not differ from subjects exposed only to clean bedding. Female rats, like ATD males, showed \(c\)-fos responses at each level of the vomeronasal pathway after being exposed to soiled bedding of sexually active males. Examples of the effect of pheromonal stimulation on Fos-IR in the mitral and granule cell layer of the AOB (peripheral level of the vomeronasal pathway) and in the mPOA (central level of the vomeronasal pathway) are shown in Figs 3 and 4. One-way ANOVA showed that exposure to soiled bedding from active males significantly augmented the number of Fos-IR nuclei in the mitral cell layer of the AOB \([F(3/13) = 10.1, P = 0.001]\) to equivalent numbers in males (ATD and cholesterol) and females (Table 2). The number of Fos-IR was also augmented in the granule cell layer of the AOB \([F(3/13) = 15.1, P < 0.001]\) in males (ATD and cholesterol) and females, with females showing a higher \(c\)-fos response than ATD and cholesterol males. Pheromonal stimulation also augmented Fos-IR in the MePV.

Fig. 4. Photomicrographs of Fos-IR in the mPOA of a male after spending 1.5 h on clean bedding (A), a cholesterol-treated male after spending 1.5 h on soiled bedding from sexually active males (B), an ATD-treated male after spending 1.5 h on soiled bedding from sexually active males (C) and a female after spending 1.5 h on soiled bedding from sexually active males (D). All animals were gonadectomized in adulthood and subsequently treated with estradiol.
At more central levels in the vomeronasal pathway, however, there were clear differences between ATD and cholesterol males in c-fos responses to odors from active males in the BNST [Kruskal–Wallis one-way ANOVA on ranks: $H = 8.5$ (d.f. = 3), $P = 0.04$] to similar numbers in both sexes. At neonatal inhibition of brain estrogen synthesis in males made their c-fos responses in the vomeronasal pathway more similar to females than control males. In peripheral portions of the vomeronasal pathway (i.e. the AOB, MePD and MePV), there were no differences between ATD and cholesterol males in the number of Fos-IR neurons. In more central portions of the vomeronasal pathway (i.e. the BNST and mPOA), differences between ATD and cholesterol males became evident. In ATD males, the BNST and mPOA were clearly responsive to odors derived from sexually active males, whereas in cholesterol males these brain areas were not responsive. The peripheral c-fos response in the cholesterol males indicates that at least they detected the male-derived odors. However, the lack of a central c-fos response indicates that the signal was differentially processed in cholesterol males as opposed to ATD males. ATD males resembled female subjects, which also showed c-fos responses at each level of the vomeronasal projection pathway after being exposed to odors of sexually active males. The behavioral findings of earlier studies predict the neuronal c-fos responses in the vomeronasal pathway when exposed to odors from sexually active males. When tested for sexual partner preference (choice: estrous female vs sexually active male), both females and neonatally ATD-treated males, gonadectomized and treated with estradiol in adulthood, spent significantly more time approaching and interacting sexually with a sexually active male than with an estrous female. By contrast, cholesterol-treated control males, castrated and estradiol-treated in adulthood, spent significantly more time approaching and interacting sexually with an estrous female than with a sexually active male. Thus, there is a sexual dimorphism in partner preference which is dependent on the neonatal action of estrogen in males. The sexual dimorphism in partner preference corresponds with the sexual dimorphism in the functional activity of the vomeronasal projection pathway, as revealed by Fos-IR, following exposure to odors from sexually active males. The sexual differentiation of the vomeronasal responses to odors of sexually active males is mediated by the neonatal action of estrogen in male subjects.

By contrast, the distribution and intensity of the increments in neuronal Fos-IR following exposure to chemosensory cues derived from the soiled bedding of estrous females was very similar in ATD and cholesterol males. Chemosensory cues derived from estrous females' bedding augmented Fos-IR in peripheral as well as in central portions of the vomeronasal projection circuit. Thus, there was no effect of neonatal estrogen deprivation on male rats' later neuronal responsiveness to odors from estrous females. From our previous behavioral findings, we expected that Fos-IR would be lowered in ATD males, because ATD males seem to be less motivated than cholesterol males to seek out the estrous female. Apparently, the behavioral findings of this previous study do not predict the neuronal c-fos responses in the chemosensory pathway when exposed to odors from estrous females. A similar discrepancy between behavior and neuronal c-fos responses was found recently by Bressler and Baum. These authors investigated odor preference (urine-soaked bedding from estrous vs anestrous females) and neuronal c-fos responses to soiled bedding from estrous and anestrous females in gonadectomized, testosterone propionate-treated male and female rats. Both males and females preferred to investigate the estrous bedding for more time than the anestrous bedding, although they had similar increments in Fos-IR after exposure to bedding from either estrous or anestrous females at each level of the vomeronasal pathway. This result was not predicted by the behavioral findings. Although partner preference is sexually differentiated, there is no clear sexual dimorphism in the functional activity of the vomeronasal system after exposure to odors from estrous females, which is surprising since large morphological sex differences exist at all levels of the pathway. The sexual differentiation of the vomeronasal pathway in males is influenced by the presence of perinatal androgens. Neonatal castration of male rats decreases the number of cells and/or volumetric measures in the vomeronasal organ, AOB, the medial posterior division of the BNST and the bed nucleus of the thalamus.
olfactory tract. Neonatal androgenization of females makes their vomeronasal pathway more male-like.31 These effects of testosterone on the developing vomeronasal projection circuit are probably mediated by its estrogenic metabolite, as has already been shown for the medial amygdala, AOB and bed nucleus of the olfactory tract.31 As shown in the present study, the vomeronasal system of ATD males functionally resembles that of females. The morphology of this system has not yet been studied in ATD males, however, except for the mPOA. In future studies, it will be important to determine whether neonatal ATD treatment blocks the masculinization of additional portions of the vomeronasal pathway (i.e. the AOB, medial amygdala and BNST) in male rats.

Neural c-fos responses to mating

Mounting and intromitting with an estrous female led to a greater neuronal c-fos expression in the mPOA of male rats, in which brain estrogen synthesis was inhibited neonatally by ATD treatment. Thus, neurons in the mPOA of ATD males, in which the normal processes of estrogen-dependent masculinization and defeminization of sexual behavior were attenuated, seem to be more responsive to mounting and intromissive stimulation than those in the mPOA of cholesterol males. The significance of the higher c-fos expression in the mPOA of neonatally ATD-treated male rats is not obvious. One possibility is that the display of masculine sexual behavior activated both stimulatory and inhibitory neurons involved in the regulation of sexual behavior in the mPOA of ATD males, whereas in cholesterol males only stimulatory neurons were activated. Supportive evidence includes the observation that ATD males first mount and intromit with an estrous female as quickly as cholesterol males. However, as the pair test progresses, the rate of mounting and intromitting decreases in ATD males, but not in cholesterol males.7 The cholesterol males continue to copulate until ejaculation is achieved. This decreased rate of mounting and intromitting in ATD males might explain why ATD males do not ejaculate as readily as cholesterol males.6,7

After eight intromissions with an estrous female, the number of Fos-IR neurons was greater in the SDN of ATD males as opposed to cholesterol males. The actual function of the SDN-POA neurons in controlling the expression of sexual behaviors is still not clear. Lesions restricted to the SDN had either no effect or a very small disruptive effect on masculine sexual behaviors in rats1,13 and ferrets,4 although in male gerbils lesions of the sexually dimorphic area strongly disrupted mating.36 The present data show that in both ATD and cholesterol males the mating-induced increments in Fos-IR occurred in portions of the mPOA which included, but were not restricted to, the SDN-POA. This is consistent with the results of lesion studies suggesting that there is no clear correlation between the function of SDN-POA neurons and the display of masculine sexual behavior in rats.1,13

CONCLUSION

The available evidence points to a sexually dimorphic processing in the central portion of the vomeronasal projection circuit of chemosensory cues derived from sexually active males as opposed to estrous females. The development of the sexually dimorphic neural response to male-derived odors results from the neonatal action of estrogen in the male rat’s nervous system.

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