Widespread accumulation of \[^{3}H\]\textit{testosterone} in the spinal cord of a wild bird with an elaborate courtship display

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ABSTRACT Elaborate courtship displays are relatively common features of the masculine reproductory behavior in birds. However, little is known about their neural and hormonal control. One bird that performs such a display is the golden-collared manakin (Manacus vitellinus) of Panamanian forests. Adult males, but not females, perform a physically intense display requiring substantial neuromuscular control of the wings and legs. We tested the hypothesis that steroid sensitivity is a property of neurons in the manakin spinal cord. Males and females were captured from active courtship leks, treated with drugs to block steroidogenesis, injected with \[^{3}H\]-labeled testosterone, and the spinal cords were removed and processed for autoradiography. Sex steroid-accumulating cells were widely distributed in the spinal cords in each of six males and in one of five females. Cells, including presumptive motoneurons, reached their highest density in the ventral horns of the cervical and lumbosacral enlargements, regions associated with motor control of the wings and legs. These results suggest that neurons in the adult manakin spinal cord can express sex-steroid receptors, but do so less in females than in males. This evidence for androgen sensitivity and sexual dimorphism in the adult avian spinal cord suggests that sex steroids may control diverse behaviors in male birds in part by acting directly on the spinal neural circuits.

To reproduce, males perform behaviors to attract and stimulate females, defend territories and mates, copulate, and care for young. Sex steroids can control the development and expression of many of these behaviors by direct actions on the central nervous system. Many studies focus on the actions of sex steroids on the hypothalamus, where there exists a relatively conserved population of neurons expressing androgen receptors (AR) and estrogen receptors (ER) within circuits controlling masculine copulatory behaviors (1–3). Steroids can also act directly on motoneurons (4, 5). For example, AR can be expressed in mammalian motoneurons of the lumbar spinal cord that innervate muscles controlling the penis (6–9). By a combination of actions of estrogens and androgens on the brain and on the spinal cord, male mammals are stimulated to copulate and are functionally able to do so.

Steroids also influence neurons controlling other reproductive behaviors (4, 10), such as the widely studied neural circuitry controlling song located in the telencephalon of the oscine passerine songbirds (11, 12). Given that song is a significant acoustic signal coordinating reproduction in these birds, it is not surprising that the song-control circuitry is influenced by sex steroids via the expression of AR and ER. But song is just one of a suite of avian reproductory behaviors. Male birds can exhibit an impressive repertoire of visual and acoustic reproductive displays (13). In some species, these visual displays can be dramatic, including acrobatic movements that are enhanced by conspicuous physical ornaments. Despite the performance of these behaviors by birds across many taxa, little is known about their hormonal and neural control. Given that some of these displays involve coordinate usage of several neuromuscular systems controlling posture and movements of the wings, legs, and tail; (ii) are often performed by males and not females; and (iii) are used in reproductive contexts, we would predict that in displaying birds, the spinal motoneurons controlling behaviorally relevant muscles would be sensitive to steroid hormones and would be anatomically and/or physiologically sexually dimorphic.

To test these hypotheses, we have performed tritiated testosterone (\(^{3}H\)-T) autoradiography on the spinal cords of adult male and female golden-collared manakins (Manacus vitellinus), a common bird species of central Panamanian forests. Manakins are a family of suboscine, passerine birds that are common in forests of the New World tropics. Males of several manakin species, including the golden-collared manakins, perform elaborate courtship displays involving short flights with midair acrobatics and intense jumping and dancing movements. In addition, the wings of some species (including golden-collared manakins) possess sexually dimorphic feather structures (14) that assist in producing loud snapping sounds by the rapid flipping of their wings (13, 14). We report that male golden-collared manakins show widespread accumulation of \(^{3}H\)-T or its metabolites in the spinal cord, including in many large motoneurons, and this pattern of sex-steroid accumulation is different in females. Androgen accumulation in spinal motoneurons in adult birds suggests that steroid sensitivity may be present in those neural pathways of the spinal cord that generate a range of avian courtship behaviors.

METHODS

Golden-collared manakins (six males; five females) were captured in mist nets from active courtship leks located in central Panama in June, 1995 and September, 1996. (All protocols for animal use have been approved by the Chancellor’s Animal Research Committee and were collected under permit from Instituto Nacional de Recursos Naturales Renovables, government of Panama.) To reduce endogenous androgen production, birds were injected immediately with an inhibitor of one of two steroidogenic enzymes, either trilostane (an inhibitor of 3-\(\beta\)-hydroxysteroid dehydrogenase/isomerase, Sterling-Winthrop Research Institute, 2 males and 1 female) or ketoconazole (an inhibitor of 17-\(\alpha\)-hydroxylase/C\(_{17-20}\) lyase, Janssen; 4 males and 4 females). Studies on the effectiveness of these two inhibitors in birds are published elsewhere (15, 16). After 24 hr (trilostane) or 8–12 hr (ketoconazole), the birds were injected with 60–80 \(\mu\)Ci (1 Ci = 37 GBq) of \(^{3}H\)-T (specific activity 102.5 Ci/mmol; New England Nuclear) and sacrificed 90 min later by decapitation. The gonads were

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Abbreviations: AR, androgen receptor; ER, estrogen receptor; \(^{3}H\)-T, \(^{3}H\)testosterone.

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RESULTS
Sex steroid-accumulating cells were found in the manakin spinal cord with substantially more cells in males than in females. Accumulation of sex steroid was found in cells of the spinal cords of all six males (on average, 209 cells over the entire cord). By contrast, only one female approached this number (with 101 cells found). Only four, two, two, and zero accumulating cells per spinal cord were found in the remaining four females.

Cells with a visible nucleus comprised 59.5% of the total number of sex steroid-accumulating cells (Fig. 1). Of the nucleated cells, 95% had a greater density of silver grains over the nucleus alone as compared with the whole soma (which includes the nucleus). Fig. 2 illustrates the total number of nucleated and nonnucleated cells with nuclear or somal accumulation (respectively) greater than 3× background. Many cells in males were present with accumulation between 3 and 5× background (64.0 on average) but were not accepted as representing significant accumulation based on our criteria. In the female in which we found a large number of cells meeting the 5× criteria, 39 additional cells met the 3× criteria. If these cells do indeed express AR or ER, but at low levels, then our results may underestimate the total number of sex steroid-accumulating cells in the manakin spinal cord (21). No additional cells that met the 3× criteria were found in the remaining females.

Sex steroid-accumulating cells with an obvious nucleus fell into two general size categories, small and large cells with mean areas of \( \approx 1,000 \mu m^2 \) and 4,000 \( \mu m^2 \), respectively (Fig. 3). Most large cells were located ventrally, whereas small cells were distributed widely, especially in middle and dorsal levels of the cord.

Even though sex steroid-accumulating cells were found in all six males, only four males provided histology that was qualitatively sufficient to allow for precise localization and mapping. Sex steroid-accumulating cells of the remaining two males were present with their size and position consistent with the large motoneurons of the ventral cervical and lumbosacral enlargements. For purposes of description, the spinal cord was subdivided rostrocaudally into the high cervical, cervical enlargement, midthoracic, and lumbosacral enlargement regions and dorsoventrally into the ventral, middle, and dorsal regions (Fig. 4). Cells were included on the map without distinguishing between those with and without nuclei. The relative abundance and distribution of sex steroid-accumulating cells in the spinal regions of the four males and five females is summarized in Table 1.

Fig. 1. Representative sex steroid-accumulating, thionin-stained neurons from the ventral cervical enlargement of an adult male golden-collared manakin.
Within the high cervical cord (rostral to the cervical enlargement), accumulating cells were found in all four males and in none of the females. Here, the number of cells were roughly evenly distributed dorsoventrally, with the middle third of the high cervical region having the most cells relative to the ventral and dorsal regions. Localization was mostly confined to an area midway between the hindbrain and the cervical enlargement (see Fig. 4), approximately C5.

In the cervical enlargement (C10–C13), sex steroid-accumulating cells were abundant in the ventral third for three of the four males and for one female. Relative to the ventral third, there were comparatively fewer cells in the middle third, and accumulation was present in all four males and none of the females. In the ventral and middle third of the cervical enlargement, cells were found along the entire rostrocaudal length. In the dorsal third, we found one male with two distinct clusters of cells, one each in the rostral and caudal portions of the cervical enlargements. We found no sex steroid-accumulating cells in this region of any other bird.

A small number of sex steroid-accumulating cells were present in the midthoracic region, approximately T3, midway between the cervical and lumbosacral enlargements. Only one male showed a few accumulating cells in the ventral third of the cord. Small cell clusters were found in two males and two females in the middle third of the cord, and a group of cells from one male was found in the dorsal third of the cord.

Sex steroid-accumulating cells were widespread in the lumbosacral enlargement (L1–L6) throughout its full rostrocaudal extent. A majority of these cells were found in the ventral third of the enlargement and were found in one male and in one female. Numerous labeled cells were also found in medial portions of the middle third of the lumbosacral enlargement, especially in two males. One of these males also had a significant number of cells in the dorsal third of the enlargement. We were unable to reliably distinguish the glycogen body on our slides.

**DISCUSSION**

These results suggest that some neurons in the manakin spinal cord express AR as described in other vertebrates (6, 9, 22). It is also possible that 3H-T was aromatized into [3H]estradiol and that ER are expressed in the manakin spinal cord, as they are in rats (23). The presence of widespread sex-steroid accumulation in cells in the spinal cords of males compared with females suggests that there is a sex difference in the magnitude of AR and/or ER expression or in the number of AR- and/or ER-expressing cells. It is also possible that sex steroid availability in the manakin spinal cord may differ between males and females because of sex differences in steroid metabolism (24, 25). Presumably, sex steroids act on spinal neural circuits in manakins to control expression of male-specific behaviors.

Many sex steroid-accumulating cells in the manakin spinal cord are located in lamina IX (20) and are of a large size, consistent with that of motoneurons (Fig. 2). Similar AR-expressing motoneurons have been found in rat (23, 26) and in *Xenopus* (27), leading us to suspect that these ventral motoneurons in manakins are also androgen-, and not estrogen-, sensitive. Some smaller sex steroid-accumulating cells found in dorsal regions of the manakin spinal cord may be sensory neurons binding estrogen as reported in rats (23) and ring doves (28). Subpopulations of AR-expressing motoneurons

<table>
<thead>
<tr>
<th>Spinal cord region</th>
<th>Bird no.</th>
<th>Ventral</th>
<th>Middle</th>
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M, male; F, female; -, no cells; +, 1–5 cells; ++, 16–50 cells; ++++, 51–100 cells; ++++, 101–200 cells; ++++++, 201–224 cells.
are functionally associated with androgen-dependent muscles and together control androgen-dependent penile reflexes in rodents (6, 25, 9) and amplexus in Xenopus (22, 27). Therefore, we assume that at least some of these sex steroid-accumulating motoneurons in manakins are also part of androgen-dependent neuromuscular systems. Because in manakins the majority of sex steroid-accumulating cells are found in the cervical and lumbosacral enlargements, and motoneurons in these enlargements largely control muscles of the upper and lower extremities (19, 29), these cells may be involved in multiple behavioral functions, perhaps innervating muscles controlling the elaborate dancing and wing-snapping of these birds.

The presence of sex differences in sex-steroid accumulation in the manakin spinal cord is consistent with the idea that sex steroids act directly on spinal neural circuits to control the expression of these courtship behaviors. Males, but not females, court actively, presumably because testosterone circulates at higher levels in males than in females. We suspect that androgens, or their metabolites, act centrally to increase the motivation to court and peripherally to increase the neuromuscular capacity to perform the displays. Androgens also act on neurons centrally and peripherally in frogs and rats to activate masculine reproductive behaviors (5, 30, 31). In the spinal cords of these species, sex differences in AR expression in part underlie the observed sex differences in behavior (5).

The basis of the sex differences in sex-steroid binding in the manakin spinal cord is unknown. The differences could arise from sex differences in the numbers of some AR-expressing cells, as observed in the rat (6, 21), in which fewer AR-expressing motoneurons exist in the female lumbar spinal cord because females lack two of the target muscles (5). It is unlikely that the female manakin possesses fewer motoneurons if they are innervating essential muscles of the wings and legs. It is more likely that higher levels of circulating testosterone in male manakins transiently up-regulate spinal AR to a greater degree than in females, as is presumed to occur in Xenopus (27). Although we cannot exclude the possibility that some of the differences we observe reflect permanent sex differences in sex-steroid binding or in other cellular attributes, we assume that neuromuscular control of the wings and legs is temporarily adapted for greater use by sex steroids when males are actively courting.

We found considerable variability in the numbers of sex steroid-accumulating cells across males and the extent to which androgens were bound by cells (Table 1). Cells were found most consistently in the cervical enlargement, but even here one male showed none. This variability in spinal cord steroid sensitivity may have been produced by differences in circulating testosterone if males were in different reproductive condition. Some birds were caught in September, a time when males are beginning to cease courtship activity (14). Our data indicate that the male with the smallest testes had the lowest number of sex steroid-accumulating cells, with the number of accumulating cells found in males with somewhat larger testes. Because androgens can directly regulate AR in the spinal cord (9), males with low plasma testosterone levels before capture could have displayed less sex-steroid accumulation. No differences in ovary size were observed that account for differences observed across females. Variability might also have been produced artificially if trilostane or ketoconazole reduced endogenous androgen synthesis differently across birds, creating disproportionate competition for androgen-binding sites with endogenous nonradioactive testosterone. We cannot exclude the possibility that different effects of these drugs on testicular or ovarian steroidogenesis might also have contributed to observed differences in spinal cord sex-steroid accumulation.

The identification of steroid-sensitive neural circuits throughout the spinal cord of the golden-collared manakin suggests that sex steroids may have a broader role in modulating avian neuromuscular systems than previously thought. Although birds have been widely studied with respect to steroid actions on the brain, no avian neuromuscular system has been fully exploited to evaluate steroid control of motoneurons and sexually dimorphic muscles that they might innervate. As in other species, sex steroids may regulate the interrelationship of motoneurons and their targets, possibly stimulating plasticity in both (32–34). Insofar as physical displays and mechanical sounds are characteristic parts of the behavioral repertoire of a vast number of bird species, the results presented here support the view that sex steroids act on the spinal cord to activate these behaviors. Further studies defining the neuromuscular control of male courtship are...
necessary to establish the role of hormones in regulating these behaviors.

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