Regulation of gonadotrophin-releasing hormone secretion by testosterone in male sheep

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In males, including the ram, testosterone, acting via its primary metabolites oestradiol and dihydrotestosterone (DHT), suppresses circulating LH concentrations. This effect is due primarily, although not totally, to decreased frequency of gonadotrophin-releasing hormone (GnRH) pulses. The arcuate-ventromedial region (ARC-VMR) of the mediobasal hypothalamus and possibly the medial preoptic area (mPOA) are sites at which oestradiol acts to suppress GnRH, but the site of DHT action is not known. Given that native GnRH neurones appear to contain few or no oestrogen or androgen receptors, the effects of testosterone metabolites probably are exerted by modulating activity of inhibitory interneurone systems such as β-endorphin, dopamine, and γ-aminobutyric acid (GABA). Although β-endorphin clearly inhibits GnRH secretion, the observation that testosterone treatment during a long-day photoperiod reduced proopiomelanocortin (POMC) mRNA in the arcuate nucleus while coincidentally suppressing GnRH release indicates that β-endorphin does not mediate the inhibitory effect of testosterone on GnRH. Activation of GABA_A receptors in either the mPOA or ARC-VMR suppressed LH, whereas activation of GABA_B receptors in the ARC-VMR increased LH pulse amplitude. Therefore, it is suggested that GABA acts in both regions to regulate LH. Whereas testosterone affects GABA metabolism in the rat hypothalamus, its effect in the ram hypothalamus is yet to be determined. Testosterone treatment activated dopaminergic cells in the retrochiasmatic A15 area in the same animals in which it suppressed POMC mRNA in the arcuate nucleus. This dopaminergic system may partially mediate the negative feedback effect of testosterone in the ram analogous to its role in partially mediating the negative effect of oestrogen in the ewe. Future studies must concentrate on determining how these and other putative inhibitory neuronal systems interact and how they in turn are regulated by environmental factors such as photoperiod.

Introduction

Upon casual observation it may be concluded that there is a relatively simple relationship among the secretory patterns of reproductive hormones in the male. A pulse of GnRH released from the hypothalamus releases a pulse of LH from the pituitary, which in turns elicits a burst of testosterone secretion from the testis. The increased testosterone then suppresses GnRH and LH, completing a typical negative feedback loop. Although it is correct, this simple depiction belies a far more complex relationship. Closer observation reveals that the feedback loop has many components and that the relationships between those components are highly dynamic, being influenced by factors such as age, photoperiod, nutritional status, and social cues. The objective of this review is to describe some of those components, how their function is modulated, and how they affect the efficacy by which testosterone regulates GnRH and LH secretion in one representative animal, the ram.
Is Metabolism of Testosterone Important?

Testosterone treatment reduces LH secretion in males of all species studied so far (Kalra and Kalra, 1989). Several observations indicate that this inhibitory effect of testosterone is mediated primarily by the testosterone metabolites oestrogen and DHT rather than testosterone per se. First, very much smaller amounts of oestrogen or DHT than testosterone are necessary for suppression of LH release (Parrott and Davies, 1979). Second, both oestrogen and DHT are produced by peripheral (Hileman et al., 1994) and neuronal aromatization and reduction (Naftolin and Ryan, 1975; Selmanoff et al., 1977) of testosterone. Third, immunization of intact rams against oestrogen greatly increases circulating concentrations of both LH and testosterone (Monet-Kuntz et al., 1988). This result could be due to neutralization of oestrogen produced by the testis as well as oestrogen produced by aromatization from testosterone. However, the observation that treatment of testosterone-treated gonadectomized rams with the aromatase inhibitor aminoglutethimide also significantly increases circulating LH (Scanbacher, 1984) provides strong evidence that oestrogen produced by aromatization from testosterone contributes significantly to normal negative feedback.

Although it was clear that administration of DHT inhibited LH release and blocked a post-castration rise in LH secretion, it was not clear whether conversion of testosterone to DHT is an essential component by which testosterone suppresses LH release. To investigate this question, we infused wethers for 72 h with either testosterone alone, reductase inhibitor alone (L-651-723 supplied by Merck Research Laboratory, Rahway, NJ), or testosterone together with reductase inhibitor (Hileman et al., 1994b). Infusion of inhibitor alone had no effect on either LH secretion or circulating concentrations of testosterone, oestrogen or DHT. In the testosterone-infused males, reduced LH release was associated with increased concentrations of all three steroid hormones. Infusion of the inhibitor with testosterone blocked only DHT formation (by over 80%) and significantly reduced, but did not completely abolish, the ability of testosterone to inhibit LH secretion (Fig. 1). These data led to the suggestion that formation of DHT is an important step in testosterone-induced reduction of pulsatile LH release. Notably, the effect of blocking both aromatase and reductase activity in testosterone-treated castrated rams has not been reported. Given the results of the cited studies, a severe or perhaps total attenuation of testosterone action on LH secretion could be expected.

Site of Testosterone Action on LH

Although the inhibitory action of testosterone on LH release is clearly established, the specific sites of action of testosterone are only partially known. Whether testosterone reduces GnRH secretion, responsiveness of the pituitary to GnRH, or both, remains unclear for some species (Kalra and Kalra, 1989). In sheep, testosterone acts primarily, although not exclusively, on the brain to suppress GnRH pulse frequency. Specifically, castration leads to increased GnRH pulse frequency (Caraty and Locatelli, 1988), whereas testosterone replacement reduces GnRH pulse frequency (Jackson et al., 1991; Tilbrook and Clarke, 1995). Circulating testosterone, at concentrations that severely reduced GnRH pulse frequency, had a marginal effect on pituitary response to exogenous GnRH (Jackson et al., 1991). However, evaluation of pituitary responsiveness across the annual breeding season leads to the conclusion that testosterone also acts directly on the pituitary to modulate the response to GnRH (Rhim et al., 1993). The neural sensitivity to the negative feedback action of testosterone as well as circulating concentrations of testosterone vary greatly with stage of the annual reproductive cycle (for example, Rhim et al., 1993). Thus it is likely that the relative effect of testosterone on the pituitary versus brain also varies with stage of the annual reproductive cycle.

The specific neural sites at which testosterone or testosterone metabolites act to regulate LH are poorly established. Both oestrogen receptor α and androgen receptor distribution in the sheep brain have been described (Lehman et al., 1993; Herbison, 1995). Within the hypothalamus there are high concentrations of oestrogen receptor α and androgen receptor in the preoptic area, arcuate and ventromedial nucleus, and median eminence. Recently, another form of the oestrogen receptor,
Fig. 1. Changes in LH pulse parameters of castrated male sheep treated with either 0.6 mg kg\(^{-1}\) 5α-reductase inhibitor (RI) L-651,723 or 768 μg kg\(^{-1}\) day\(^{-1}\) testosterone (T), or T + RI for 3 days. IPI: interpulse interval. Bars represent means ± SEM of day 3 values minus day 0 values (n = 5 per treatment). P values are indicated within each graph for comparisons made between groups using tests of least significant differences. (Reproduced from Hileman et al., 1994 with permission.)

Oestrogen receptor \(\beta\) has been found. Although the function of oestrogen receptor \(\beta\) is not clear, its distribution in the male sheep hypothalamus is similar to that described for the rodent (Shugrue et al., 1997) with localization in the medial preoptic area, retrochiasmatic area, bed nucleus stria terminalis, paraventricular nucleus, supraoptic nucleus and dorsomedial hypothalamus. Only sparse labelling for oestrogen receptor \(\beta\) is found in the arcuate nucleus and ventromedial hypothalamus (S. M. Hileman and R. J. Handa, unpublished). Thus, each of these sites, as well as others in the brainstem and amygdala, may be involved in mediating the action of testosterone on GnRH secretion.

In an attempt to delineate specific sites at which testosterone, oestrogen, and DHT act to
suppress LH, we (Scott et al., 1997) placed implants of these steroids into the mPOA and ARC–VMR of the ventromedial hypothalamus of long-term castrated rams. Implants of testosterone and DHT at either site were ineffective at suppressing LH. In contrast, implants of oestrogen in the mPOA were marginally effective whereas implants of oestrogen into the ARC–VMR clearly suppressed LH (Fig. 2). These results implicate the ARC–VMR as an important site at which oestrogen acts to suppress LH secretion. The reason for the failure of testosterone and DHT implants to suppress LH is not clear, but may reflect less efficient diffusion, downregulation of androgen receptors following castration (Handa et al., 1996), or the fact that androgens may act at other or additional untested sites.

By Which Neural Pathways Does Testosterone Act?

The specific mechanisms by which testosterone or its metabolites act to suppress GnRH release are not clear. The most parsimonious mechanism would be action of the steroids directly on GnRH neurones. Recent observations indicating that immortalized GT1-7 GnRH-secreting cells contain both androgen and oestrogen receptors support this contention (Belsham et al., 1998; Shen et al., 1998). However, it is debatable whether these cells are fully representative of endogenous GnRH secreting cells in vivo. In addition, it is notable that several investigations have found few or no steroid receptors on native GnRH cells in vivo (Shivers et al., 1983; Huang and Harlan, 1993; Lehman and Karsch, 1993). Consequently, the prevailing hypothesis is that steroids affect GnRH neurones through actions on interneurones. This idea gains support by observations that numerous neurotransmitter agonists and antagonists affect GnRH secretion and that several neuronal systems concentrate gonadal steroids. Under this concept, steroids could reduce GnRH secretion either by increasing secretion of inhibitory neurotransmitters or by reducing secretion of stimulatory neurotransmitters or by a combination of effects. Owing to the inherent difficulty of such studies, relatively little effort has been given to investigating the second or third possibilities. The first possibility is perhaps the easiest to address and results of several studies lead to the suggestion that gonadal steroids modulate the secretion of at least three neurotransmitters known to inhibit LH secretion: opiates (i.e. β-endorphin), dopamine, and γ-aminobutyric acid (GABA).

β-Endorphin

β-Endorphin neurones are found in high concentrations in the arcuate nucleus, contain oestrogen receptors, and contact GnRH neurones (Leranth et al., 1988; Thind and Goldsmith, 1988; Barb et al., 1991; Lehman and Karsch, 1993). Numerous data show that administration of the opiate agonist morphine suppresses LH, whereas injection of antagonists such as naloxone increases LH (see Barb et al., 1991). Observations that the effects of opiate antagonists are much more robust in intact or testosterone-treated than in castrated animals led to the concept that opiates may mediate the inhibitory action of testosterone (Ebling and Lincoln, 1985; Barb et al., 1991). The observation that peripheral concentrations of β-endorphin were highest during the breeding season (Ssewannyana and Lincoln, 1990) indicates that the influence of endorphin is greatest during that period. Intuitively, this seems inconsistent with the fact that testosterone is relatively ineffective at inhibiting GnRH release at this time. To investigate this issue further, we performed a series of experiments (Hileman et al., 1996; Hileman et al., 1998) to examine the effect of testosterone on POMC mRNA in the arcuate nucleus under the assumption that amounts of the precursor mRNA may reflect the synthesis and secretion of the peptide. In the most recent study, males were either castrated or castrated and implanted with testosterone, and then placed under either an inhibitory long-day or stimulatory short-day photoperiod. Testosterone did not alter either LH release or amounts of POMC mRNA in animals exposed to short days. In contrast, testosterone greatly reduced both mean LH concentrations and the amount of POMC mRNA (Fig. 3), but not of GnRH mRNA, in animals exposed to long days (Fig. 3). This finding was consistent with our previous report (Hileman et al., 1996) that testosterone administered for either 3 days or 3 months reduced POMC mRNA in the
Fig. 2. Examples of LH secretory profiles in plasma of three castrated rams with either stylets or steroid implants placed bilaterally into a site located dorsal-lateral to the arcuate nucleus and medial to the ventromedial nucleus of the hypothalamus (VMH). CON: stylet; CHOL: cholesterol; DHT: dihydrotestosterone; E: oestradiol. Peaks of LH pulses are represented by hollow circles. Sampling periods were 7 days apart. The number in the upper right of each panel refers to an individual animal identification number. (Modified from Scott et al., 1997.)
arcuate nucleus of wethers kept under ambient long days. Although it is recognized that mRNA content may not always reflect actual peptide release, studies from two other laboratories independently investigating the effects of feed restriction on POMC mRNA content and β-endorphin release indicate that relative steady-state amounts of POMC mRNA are indicative of relative hypothalamic β-endorphin release rates in the lateral median eminence of sheep (Prasad et al., 1993; McShane et al., 1993). In addition, preliminary data from our recent studies using portal-cannulated males indicate that naloxone effectively increases GnRH and LH release in castrated males exposed to inhibitory long days. This finding suggests that the magnitude of response to naloxone may be dependent more on the basal LH secretory rate than on steroid background. These findings support the concept that the increased synthesis of β-endorphin is not a mechanism whereby testosterone suppresses GnRH release in animals exposed to long days. It is noteworthy that Goodman et al. (1995) reached a similar conclusion regarding the relationship between oestrogen, the opiates, and LH secretion in ewes.

Dopamine

Dopaminergic neurones are clumped into nuclei or groups scattered throughout the brain. Small groups designated A12, A14, A15, and located in the anterior part of the hypothalamus appear to have a significant role in regulating LH in sheep. The A14–A15 group is located in the retrochiasmatic area; the A12 group is located in the arcuate nucleus–median eminence region. A large body of evidence from studies using female sheep and a variety of approaches led to the concept that dopaminergic input from these areas inhibits LH secretion, at least when the animals are exposed to long-day photoperiods (Thiery et al., 1995; Lehman et al., 1996; Viguie et al., 1997).

In comparison, there are relatively few studies on the role of the dopaminergic system on LH secretion in rams, but most of these data support an inhibitory role. Although pimozide, a dopamine

Fig. 3. Effects of testosterone (T) and duration of photoperiod on silver grain area per neurone (solid bars) and numbers of proopiomelanocortin (POMC) mRNA positive cells (■) in the arcuate nucleus of castrated rams. LD: long days (16L:8D); SD: short days (10L:14D). Data are presented as means ± SEM. POMC mRNA grain area in the LD + T group was lower \((P < 0.01)\) than in the other groups which did not differ from each other \((P > 0.10)\). There was an effect of steroid \((P < 0.05)\) and a strong tendency for an effect of photoperiod \((P = 0.06)\), but no interaction of treatments \((P > 0.40)\), on the number of POMC cells. (Reproduced from Hileman et al., 1998.)
Testosterone and LH

D, receptor antagonist, did not increase LH in rams (Tilbrook and Clarke, 1992), the more specific D, antagonist sulpiride was effective in Soay rams, particularly during the non-breeding season (Tortoneze and Lincoln, 1994). In addition, supporting evidence was obtained in our laboratory by monitoring the effect of testosterone administration on expression of the early-intermediate gene c-Fos in the A14-A15 cell groups. We used dual-label immunocytochemistry for c-Fos and tyrosine hydroxylase (the rate-limiting enzyme in dopamine formation) to determine the percentage of dopaminergic cells activated by testosterone. The tissue used was from our previously cited study in which we had shown that infusion of testosterone for 3 days suppressed LH release coincident with reduced POMC mRNA (Hileman et al., 1996). Testosterone treatment significantly increased the percentage of dual-labelled cells in the A15 group and strongly tended (P < 0.06) to increase that percentage in the A14 group. No effect of testosterone was noted on either the A13 cell group or the total number of tyrosine hydroxylase positive cells (Lubbers et al., 1995). These results are very similar to those obtained from ewes treated with oestrogen during the non-breeding season (Lehman et al., 1996). In summary, there is evidence to support the concepts (1) that activation of the A14-A15 cell groups is involved in the feedback action of gonadal steroids during the non-breeding season and (2) that during this season gonadal steroids selectively activate this dopaminergic subsystem in both rams and ewes.

GABA

Gamma-aminobutyric acid (GABA) is a widely distributed neurotransmitter, the primary action of which is to inhibit the activation of other neuronal systems. Several observations indicate that it acts in the mPOA of rats to inhibit GnRH release, and results of experiments in male rats support the concept that GABA may mediate the inhibitory effect of testosterone on GnRH and LH release, particularly within the mPOA (Grattan and Selmanoff, 1993; Grattan et al., 1996; Sagrillo and Selmanoff, 1997). Results of studies in ewes also led to the contention that GABA acts in the mPOA to suppress GnRH secretion (Robinson, 1995; Scott and Clarke, 1993a,b).

Work from our laboratory supports the concept that GABA acts in both the mPOA and ARC-VMR to suppress GnRH release and within the ARC-VMR to regulate specifically LH pulse amplitude (Ferreira et al., 1996). Castrated rams had guide tubes stereotaxically placed bilaterally either in the mPOA or ARC-VMR to deliver drugs into these specific sites by microdialysis. Subsequently, these areas were perfused with artificial cerebrospinal fluid for 4 h followed by 4 h of either cerebrospinal fluid, the GABA_A receptor agonist muscimol, or the GABA_B receptor agonist baclofen. In the mPOA, muscimol treatment reduced pulsatile LH release, whereas baclofen was without effect. In the ARC-VMR, muscimol also inhibited pulsatile LH release, but surprisingly baclofen actually increased LH release. This effect was due primarily to an increase in pulse amplitude rather than pulse frequency and probably reflected an increase in GnRH pulse amplitude (Fig. 4). Thus, it appears that GABA may act through the GABA_A receptor to suppress pulsatile GnRH release in both the POA and ARC-VMR. Interpreting the effect of baclofen is not as straightforward, but may be explained by the fact that GABA_B receptors apparently function as autoreceptors. Local stimulation of presynaptic autoreceptors may reduce local secretion of endogenous GABA and thus free GnRH neurones from chronic inhibition.

Although activation of the GABA_A and GABA_B receptor subtypes obviously altered LH release, results from our studies addressing the issue of whether these receptor types mediate testosterone negative feedback are more difficult to interpret. The GABA_A receptor antagonist bicuculline methiodide (BMI) and the GABA_B receptor antagonist CGP 55854A were administered during the breeding season into only the ARC-VMR of castrated males and castrated testosterone-treated males. The expectation was that the GABA_A antagonist would increase LH. However, BMI consistently suppressed LH release in castrated males and failed to increase LH secretion in the testosterone-treated males. CGP 55854A was without effect in either group. The apparently paradoxical effects of BMI might be explained by inhibitory effects of this drug on N-methyl-D-aspartic acid (NMDA) receptors or other neurotransmitters (Svenneby and Roberts, 1973; Miller and
Fig. 4. Plasma LH profiles of one castrated ram subjected to separate sequential bilateral microdialysis infusion of artificial cerebrospinal fluid (aCSF) only (a), or aCSF followed by 1 mmol baclofen \(1^{-1}\) (b), or aCSF (c) followed by 1 mmol muscimol \(1^{-1}\) into the arcuate–ventromedial region of the hypothalamus. The drug concentrations listed are those of the dialysis solution. It is estimated that the total doses of baclofen and muscimol delivered at each site were 7.9 and 4.5 pg, respectively. Note the enlarged scale of the middle panel. Peaks of LH pulses are represented by hollow circles. Drug delivery started at 4 h, as indicated by the vertical line. (Reproduced from Ferreira et al., 1996.)

McLennan, 1974; Krebs et al., 1994; Musshoff et al., 1994) and in retrospect indicates that studies using this antagonist are interpreted with caution.

However, it should be noted that although BMI injected into the POA of ovariectomized ewes consistently reduced LH when given during the breeding season, it increased LH in some oestrogen-treated ovariectomized ewes when given during the anoestrous period (Scott and Clarke, 1993b). Thus, the effect of BMI on LH may vary with steroidal background or season of treatment. Possibly, during the breeding season the GABAergic system is relatively inactive, particularly in the absence of gonadal steroids, and the only observable effect of BMI on LH secretion is inhibition due to blockade of other essential stimulatory systems attained by relatively high doses of the drug. In contrast, during the non-breeding season, when GABA activity is postulated to be increased by...
Fig. 5. Schematic illustration of the possible mechanisms by which testosterone (T) controls release of gonadotrophin-releasing hormone (GnRH) in male sheep. (+): stimulatory action; (−): inhibitory action; E: oestradiol, DHT: dihydrotestosterone; β-end: β-endorphin secreting neurone; DA: dopamine secreting neurone; GABA: γ-aminobutyric acid secreting neurone; A: GABA_A receptor.

gonadal steroids, the GABAergic system may become more sensitive to blockade. Although it is clear that GABA modulates LH secretion in rams, data are insufficient to conclude whether it partially mediates the action of testosterone on LH. Clearly, additional study will be required to resolve these issues.

Summary Model

Results of studies from several laboratories are incorporated into a summary model (Fig. 5) illustrating some of the possible neurochemical pathways by which testosterone alters GnRH release in rams. The first point is that the effects of testosterone are mediated largely, if not exclusively, by the metabolites oestrogen and DHT. These steroids act in the ARC—VMR and probably other sites to reduce GnRH pulse frequency, but not GnRH synthesis. A second point is that these steroids probably do not act directly on GnRH neurones, but act indirectly by modulating secretion of one or more inhibitory neuromodulators such as β-endorphin, dopamine and GABA.

The specific role of β-endorphin remains unclear, but there is overwhelming evidence that it acts tonically to inhibit GnRH release. On the other hand, it does not appear to mediate the inhibitory action of testosterone. Our results support the postulate that testosterone suppresses, not stimulates, synthesis of β-endorphin, particularly when the animals are exposed to a long-day photoperiod. It appears paradoxical that when testosterone is maximally suppressing GnRH release, it also is suppressing a system that inhibits GnRH release. However, the model deals with this by incorporating the possibility that testosterone also increases local dopamine release, particularly during the non-breeding season. Stimulation of dopamine release probably has two outcomes. First, DA may act directly to suppress GnRH release. Second, DA also may act to inhibit β-endorphin secretion (Tortonese and Lincoln, 1994). In our study in which testosterone suppressed POMC mRNA (Hileman et al., 1996), it also activated A14 and A15 dopamine neurones (Lubbers et al., 1995). As β-endorphin neurones contain oestrogen receptors, there may also be a direct effect of this testosterone metabolite. However, it is not clear that these associated changes reflect cause and
effect. As indicated in the model testosterone-induced reduction in β-endorphin secretion could also secondarily lead to increased dopaminergic activity. In either case, if the inhibitory effect of dopamine on GnRH secretion is relatively stronger than that of β-endorphin, activation of this network by testosterone will still suppress GnRH release. In addition, it should be noted that the relationship among testosterone and the neurotransmitters is not static. During the breeding season the effectiveness of testosterone in altering the activity of these systems and in suppressing GnRH is reduced. Although the suppression of β-endorphin may be reduced, resulting in greater β-endorphin release, a parallel reduction in dopamine release would ultimately result in increased GnRH release.

In the model we suggest that GABA inhibits GnRH release by acting via GABA$_A$ receptors in both the mPOA and ARC-VMR. However, the indication that GABA partially mediates the action of testosterone is speculative.

Conclusions

There are several important unanswered questions about how testosterone reduces GnRH and LH secretion. Although oestrogen and DHT appear to mediate the action of testosterone, the relative contribution of these two metabolites is unclear. Suppression of either DHT or oestrogen formation partially blocked the inhibitory effects of testosterone on pulsatile LH release. The effect of simultaneously blocking both reductase and aromatase activities has not been determined. In addition, it is not clear exactly where in the hypothalamus or brainstem oestrogen and DHT act to alter GnRH release. Indeed, it is yet to be clearly demonstrated that DHT alters GnRH release. This issue requires further investigation, perhaps using a more androgen-responsive model than the long-term castrated male.

The specific role of the various neuromodulators remains unresolved. We are currently addressing the question of whether testosterone stimulates GABA release in the hypothalamus. However, the specific physiological roles of the GABA receptor subtypes, and the respective roles of the widely distributed GABA neurones in modulating or mediating the effects of testosterone on GnRH release remain to be elucidated fully in the ram.

The specific role of β-endorphin, and of other opiates, also remains elusive. It is unknown whether testosterone alters hypothalamic β-endorphin release and coincidentally whether hypothalamic release of β-endorphin reflects changes in steady state POMC mRNA. Furthermore, it is not known whether testosterone or environmental factors regulate opiate receptors, nor is it clear exactly where the opiates act. The report by SaneIla et al. (1997) that GnRH neurones lack opiate receptors leads to the suggestion that still other interneurone systems must be involved – perhaps those secreting nitric oxide (Brann and Mahesh, 1997; Lopez et al., 1997).

Dopamine neurones of the A14 and A15 groups appear to be activated by testosterone during long days. Whether this is specific for an inhibitory photoperiod in males as in females is not known. The relative importance and relationship of this subset, and the A12, dopamine neurones to GnRH release has not been determined nor have efferent and afferent pathways to and from these dopamine groups. Given that the A14–A15 dopaminergic neurones apparently contain little oestrogen receptor-α (Lehman and Karsch, 1993) it is not clear how their function is modulated by either photoperiod or steroids.

In addition, there is a fundamental question as to how photoperiod acts to gate the sensitivity of these and possibly other systems to testosterone. Ultimately this involves melatonin, the pineal hormone which transduces photic information into a chemical signal. Although considerable progress has been made (for example Malpaux et al., 1998; Hileman et al., 1994a) neither the neuroanatomical components nor the identity of neural systems involved in this pathway are completely known. Perhaps even more perplexing is the consistent observation that although testosterone or oestrogen administration suppresses LH pulse frequency, pharmacological manipulations of various neurotransmitters have failed to mimic fully the action of either steroid treatment or removal. Consequently this leads to the suggestion that testosterone, its metabolites or
both may act through the coordinated activity of several neurotransmitter systems rather than a single system. Thus, it seems likely that much effort will be required before we fully understand the ‘relatively simple’ pathways by which testosterone regulates GnRH and LH release in males before those by which environmental factors modulate the action of testosterone can be determined.

The authors gratefully recognize the contributions of our colleagues Suzie Ferreira, David Kuehl, Laura Lubbers, and Chris Scott who contributed to the work from this laboratory. These studies were supported by NIH Grant HD 27453 and USDA Grants AG92-37202-8177 and AG95-37203-2033.

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Testosterone and LH 241
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