Histomorphology, oLH and hCG receptors, and testosterone secretion in vitro in Rambouillet rams from lines in which females had been selected for low or high reproductive rate

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The objective of this study was to determine whether gross or histomorphological components of the testes, capacity and dissociation constants (Kd) of testicular oLH and hCG receptors, and gonadotrophin-stimulated testosterone secretion in vitro differed among Rambouillet rams from lines selected for low or high female reproductive rate and from rams of a random-bred control line. Lines had been selected for approximately 20 years. Data were collected from 22-month-old rams during the late breeding season. Rams among lines did not differ (P > 0.05) in gross testicular characteristics or most histomorphological characteristics. However, the percentage volume of interstitial vascular tissue was greater (P < 0.05) for rams from lines selected for low female reproductive rate than for rams from lines selected for high female reproductive rate. Receptor sites per Leydig cell and binding capacities of oLH and hCG receptors per testis, per gram of parenchyma, and per milligram of membrane protein did not differ (P > 0.05) among lines. The Kd values for oLH and hCG receptors did not differ (P > 0.05) among lines; however, receptor sites per Leydig cell, capacities of testicular parenchyma to bind gonadotrophin and Kd values were higher (P < 0.05) for oLH than for hCG receptors. Total oLH- and hCG-stimulated testosterone secretion in vitro did not differ (P < 0.05) among lines. In conclusion, selection for or against reproductive rate in Rambouillet ewes has not altered gross or most histomorphometric characteristics of the testes of male offspring, with the exception that selection against reproductive rate increased the proportion of testicular volume occupied by vascular tissue within the interstitium. Furthermore, selection has not altered total oLH- and hCG-stimulated testosterone secretion in vitro or the affinity and capacity of oLH and hCG receptors of the testes of male offspring.

Introduction

Breeds of sheep that differ in reproductive rate display positive correlations between traits of female reproductive rate and certain male reproductive characteristics such as testicular diameter (Land, 1973; Knight, 1984) or serum concentrations of LH (Thimonier and Pelletier, 1972; Land and Carr, 1975; Carr and Land, 1975). On the basis of such relationships, it was hypothesized (Bindon and Turner, 1974; Land, 1974) that certain aspects of male and female reproduction may be controlled by the same physiological factors that in turn are controlled by genes common to males and females. Chubb (1992) reported that genes controlling testis size are not directly associated with the Y chromosome. Therefore, it is possible that selection based upon a sex-limited characteristic of either sex would produce changes in genetically correlated characteristics of the opposite sex.

Data that support this hypothesis are as yet equivocal for females from lines in which males had been selected for or against reproductive endocrine responses or testicular traits (Haley et al., 1989, 1990; Matos and Thomas, 1992). Studies that evaluated reproductive endocrine responses and certain testicular characteristics of rams from lines of sheep in which females had been selected for different reproductive rates indicated little or no correlated responses in male offspring (Bindon et al., 1985; Hochereau-de Riviers et al., 1990a; Curry et al., 1993).

Most studies in this area have focused upon the relationship between characteristics of female reproductive rate as the criterion for selection and male reproductive traits such as LH or FSH concentrations, scrotal circumference, testicular diameter, and testicular mass. However, the effects of selection for female reproductive rate upon histomorphological and endocrinological characteristics of the testis have not been well documented.

Two lines of Rambouillet sheep that differ in reproductive rate were established by applying selection to a female sex-limited trait (Schoenian and Burfening, 1990). The selection

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process for establishing these lines has induced a change in the ovulation rate without affecting embryonic mortality (Schoenian and Burfening, 1990). The fact that an ovarian trait has been altered by selection led us to hypothesize that perhaps selection within these lines had induced correlated changes in testicular morphology and physiology in rams of these lines. The objectives of this study were (i) to determine whether selection for or against female reproductive rate in Rambouillet sheep had altered scrotal circumference, testis mass, testis tone and histomorphological components of the testicular parenchyma, and (ii) whether there were differences in histological components, to determine whether they were reflected in changes in either the capacity or affinity of testicular oLH and hCG receptors, or the ability of testicular parenchyma to secrete testosterone in vitro in response to gonadotrophin of mature male offspring.

Materials and Methods

Animals and trials

Rambouillet rams from lines in which ewes had been selected for either high (HL) or low (LL) reproductive rates, and Rambouillet rams from a random-bred control line (CL) were used in this study. Two trials were performed in consecutive years. Lines had been selected for female reproductive rate for 19 (Trial 1) and 20 (Trial 2) years. The details of the establishment, selection procedure, nutritional management and handling of these lines have been described by Schoenian and Burfening (1990). Each trial began during the first week of December and ended during the last week of February. At the start of each trial rams were 22 months of age and rams from each line were allocated randomly into three blocks to be orchidectomized at the beginning (first 3 weeks), middle (second 3 weeks), and end (third 3 weeks) of each trial. This method was used as a means to evaluate whether time (season) interacted with the type of line to bias the results of this experiment.

Scrotal circumference, tone and orchidectomy

The scrotum of each ram was shaved and the testes pushed towards the bottom of the scrotum to measure scrotal circumference at its widest point. After measuring the scrotal circumference, the pressure on the testes was released and the tone of each testis was determined by applying a tonometer (Lane Mfg Inc., Denver, CO) to the external scrotal surface of the medial anterior area of each testis. Immediately thereafter, each ram was bilaterally orchidectomized under a surgical plane of anaesthesia induced and maintained with i.v. infusion of thiamylal sodium into one jugular vein (4% in 0.9% sterile saline, Bio-CEutics Laboratories, Inc., St Joseph, MO). In Trial 1, the mass of each testis was recorded after removing the epididymis and tunica albuginea. In Trial 2, only the mass of the left testis was obtained because the right testis was prepared for perfusion fixation to evaluate histomorphological characteristics.

Testis perfusion and slide preparation

In Trial 2, the right testis of each ram was perfused immediately after orchidectomy by a procedure modified from Glaubert (1975). Perfusions were performed at 22°C through the testicular artery at a point close to the caput epididymis with 3% glutaraldehyde:1% formaldehyde solution in 0.075 mol cacodylate buffer 1-1. The perfused testis was cut transversely into three parts: top, middle and bottom. Four 2 mm x 2 mm x 4 mm pieces were cut from each part and fixed overnight at 4°C by immersion in buffered 2% glutaraldehyde. The pieces were then postfixed in buffered 1% OsO4 for at least 2 h, dehydrated in ethanol and embedded in araldite epoxy resin 502. Five thin sections (1 µm) were cut from each block of embedded tissue and stained with Toluidine blue and basic fuchsin as described by Hoffmann et al. (1983). Two of the five sections from each block were chosen at random and photographed at x 125 magnification using a light microscope. Details of two representative areas of the interstitium within each area included in the photomicrographs taken at x 125 were then photographed at x 1250 magnification.

Stereology

The total areas occupied by seminiferous tubules, interstitium and cross-sectional areas of seminiferous tubules were determined in the photomicrographs taken at x 125. These measurements were then used to estimate the proportions of the volume of the testis (volume percentages) occupied by seminiferous tubules and interstitium, as described by Lunstra and Schanbacher (1988), and seminiferous tubule diameters (A = πr²), respectively.

Areas occupied by Leydig cells with and without nuclei, vascular tissue and all other structures of the interstitium collectively were measured in the photomicrographs taken at x 1250. These determinations were used to estimate the volume percentages of each of these components of the interstitium and for the whole testis (Lunstra and Schanbacher, 1986). Cross-sectional areas of Leydig cell nuclei were also determined in the photomicrographs taken at x 1250. Areas were measured directly from photomicrographs using a digitizer pad (Kurta® IS/ONE, Phoenix, AZ) and calculations were performed by a computer using Sigma-Scan® (Jandel Scientific, Corte Madera, CA).

Leydig cells and Leydig cell nuclei were assumed to be round and their volumes were obtained by first calculating the diameter from the cross-sectional area (A = πr²), adjusting the diameter according to Abercrombie (1946), and then using this corrected value to calculate the volume (V = 4πr³/3).

The number of Leydig cells per gram of parenchyma was calculated by dividing the total volume of Leydig cells in 1 g of parenchyma by the Leydig cell volume (Allrich et al., 1983). A similar procedure was used to calculate the number of Leydig cells per testis. The specific gravity of the testicular parenchyma of each ram was determined and the average value of 1.04 was used for all calculations of testicular volume because a preliminary analysis demonstrated that specific gravity did not differ (P > 0.05) among lines.

Testosterone secretion in vitro

Immediately after orchidectomy, the tunica albuginea of the left testis was removed and one longitudinal half of the
parenchyma was processed for evaluation of testosterone secretion in vitro. The other half of the parenchyma was frozen at \(-20^\circ\)C for the later assay of oLH and hCG receptors. Testicular parenchyma was processed for stimulation of testosterone secretion in vitro using the procedure of Berardinelli et al. (1989). Incubations were performed under an atmosphere of 5\% CO\(_2\)–95\% O\(_2\) for 4 h. Tissue was placed into flasks containing ME 199 medium (Cellgro; Mediatech, Washington, DC). The ME 199 medium in each flask was discarded and replaced three times with fresh medium at 15-min intervals before adding the gonadotrophins. The time from orchidec-tomy to the time when tissue rinsing began was approximately 40 min. Tissue rinsing was necessary because it removed residual testosterone and probably gonadotrophins present in the fluid accompanying the tissue mince that could interfere with gonadotrophin-stimulated testosterone secretion in vitro and its evaluation (Berardinelli et al., 1989).

**Gonadotrophins and dosages**

hCG was used to stimulate testicular tissue in Trial 1 and both hCG and oLH were used in Trial 2. Ten minutes after tissue rinsing, hCG (US Biochemical Corporation, Cleveland, OH) or oLH (NIDDK-oLH-I-3, Bethesda, MD) was added to flasks containing tissue in ME 199. Dosages of hCG were 0, 0.025, 0.25 and 2.5 µIU ME 199 ml\(^{-1}\). Dosages of oLH were 0, 0.13, 1.3 and 130 ng ml\(^{-1}\). Dosages of each gonadotrophin were prepared in ME 199 immediately before use, evaluated in quadruplicate, and added to each flask in 100 µl aliquots. Dosages of oLH were equivalent on a molecular basis to dosages of hCG. Equivalencies were calculated taking the molecular mass of oLH as 28.3 kDa and that of hCG as 37 kDa (Sairam et al., 1988) and the equivalence of 15 000 IU of hCG per mg of the hormone. Thus the concentrations of gonado-trophins tested were 0, 4.6, 46 and 4600 pmol l\(^{-1}\).

**Sampling and testosterone radioimmunoassay**

Samples (300 µl) of ME 199 were taken from each flask just before addition of gonadotrophin (0 h) and after 1, 2, 3 and 4 h of incubation for testosterone determination. Testosterone was assayed without extraction by a radioimmunoassay validated in our laboratory (Byerley et al., 1990). Assay sensitivity was 20 pg ml\(^{-1}\). Interassay coefficients of variation for pools of samples that inhibited binding at 29 and 75\% were 4.5 and 9.2\%, respectively, and intra-assay coefficients of variation for these pools were 4.9 and 9.9\%, respectively.

**oLH and hCG receptor assays**

Iodination grade oLH (NIDDK-oLH-I-3, Bethesda, MD) and hCG (hCG no. CR-125, NICHID, NIH, Bethesda, MD) were labelled with \(^{125}\)I (New England Nuclear, Boston, MA) by the lactoperoxidase method and purified by gel filtration chromatography in Sephadex G-75 columns (Pharmacia, Uppsala). Hormones were eluted from columns with 25 mmol Tris–HCl 1\(\times\) (pH 7.5) containing 10 mmol MgCl\(_2\) 1\(\times\) and 1 mg BSA ml\(^{-1}\). Elution fractions that had at least 70\% of the total radioactivity in the TCA precipitate were used for the assays. Specific activities of each hormone, based upon the percentage of incorporation in TCA precipitants, were 27.4 µCi µg\(^{-1}\) and 38.9 µCi µg\(^{-1}\) for \(^{125}\)I-labelled oLH and \(^{125}\)I-labelled hCG, respectively.

Testicular membranes for radioreceptor assays were prepared by a modified procedure from Sairam et al. (1988). Briefly, frozen testicular parenchyma was thawed overnight at 4°C. All further processing of membrane preparations was performed at this temperature. Approximately 25 g of testicular parenchyma was weighed, cut into small pieces with scissors, homogenized using a homogenizer (Brinkmann Instruments Co.), and then filtered through four layers of gauze. Homogenizations were carried out in 50 ml of Tris–HCl buffer (25 mmol l\(^{-1}\), pH 7.5) containing 25 mmol sucrose l\(^{-1}\). The filtrate was centrifuged initially at 500 g for 15 min, the supernatant was then recovered and centrifuged at 30 000 g for 30 min. The pellet was resuspended in Tris–HCl buffer (25 mmol l\(^{-1}\), pH 7.5) and centrifuged again at the same speed. Cen trifugations were carried out at 4°C. Pellets were resus- pended in Tris–HCl buffer (25 mmol l\(^{-1}\) containing 10 mmol MgCl\(_2\) 1\(\times\)). Protein concentrations of membrane suspensions were determined with the BCA\(^\text{®}\) Protein Assay (Pierce, Rockford, IL), using BSA as the standard. Increasing amounts (0.0625, 0.125, 0.25, 0.5, 1, 2, 4, 8, 16 and 24 ng) of \(^{125}\)I-labelled oLH or \(^{125}\)I-labelled hCG were incubated in triplicate for 22 h at 4°C with testicular membranes equivalent to 1.5 mg of protein and assay buffer to a final volume of 500 µl per tube. The assay buffer was Tris–HCl (25 mmol l\(^{-1}\), pH 7.5) containing 10 mmol MgCl\(_2\) 1\(\times\) and 1 mg BSA ml\(^{-1}\). Nonspecific binding was determined in the presence of 1 µg of unlabelled hCG or oLH per tube, respectively. The binding of hormone and receptor was stopped by adding 2 ml of cold assay buffer followed by centrifugation at 3000 g for 20 min at 4°C. Radioactivity was determined in the pellets (counts bound) and supernatants (counts free) in a Packard 5160 auto-γ scintillation spectrometer with an efficiency of 58.2\%.

The maximum binding capacity of labelled hormones used in the assays was assessed by incubating 0.05–0.5 ng of each hormone with an excess of sheep testicular membrane preparations and were found to be 17.2\% for \(^{125}\)I-labelled oLH and 21.3\% for \(^{125}\)I-labelled hCG. All binding assays were performed with a single preparation of each labelled hormone.

**Receptor binding capacity and K\(_d\) values**

Receptor binding capacity and the dissociation constant for each hormone were estimated by Scatchard plot using the computer program Enzfitter (Elsevier-BIOSOFT, Cambridge, UK). For the purpose of these analyses, the amount of radioactivity specifically bound was obtained by subtracting the nonspecific binding counts from the counts bound (counts in the pellets).

Specifically bound radioactivity was transformed into ng of hormone using the specific activity and the equivalence of 1 µCi = 2.2 × 10\(^{8}\) d.p.m. The amount of free hormone was calculated by subtracting the amount of hormone specifically bound from the total amount of hormone added to the assay tube. Molar concentrations of free and bound hormone were estimated using the molecular masses for oLH and hCG given above.
Table 1. Least-squares means for testis volume (TV) and percentage volumes of seminiferous tubules (ST), interstitium (INTS), vascular tissue (VT), Leydig cells (LC) and other components of the testicular interstitium of Rambouillet rams from lines selected for low (LL) or high (HL) reproductive rates and from a random-bred control line (CL).

<table>
<thead>
<tr>
<th>Selection line</th>
<th>n</th>
<th>TV (cm^3)</th>
<th>ST (%)</th>
<th>INTS (%)</th>
<th>VT (%)</th>
<th>LC (%)</th>
<th>Other (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LL</td>
<td>8</td>
<td>91.0</td>
<td>80.7</td>
<td>19.3</td>
<td>4.6*</td>
<td>2.3</td>
<td>12.5</td>
</tr>
<tr>
<td>HL</td>
<td>6</td>
<td>85.2</td>
<td>81.6</td>
<td>18.4</td>
<td>3.6b</td>
<td>2.7</td>
<td>12.0</td>
</tr>
<tr>
<td>CL</td>
<td>7</td>
<td>86.5</td>
<td>82.2</td>
<td>17.8</td>
<td>3.6b</td>
<td>2.2</td>
<td>11.8</td>
</tr>
<tr>
<td>SEM</td>
<td></td>
<td>24.8</td>
<td>3.4</td>
<td>3.4</td>
<td>0.8</td>
<td>0.5</td>
<td>2.7</td>
</tr>
</tbody>
</table>

*Values within columns with different superscripts are significantly different (P < 0.05).
*Pooled SEM.

Statistical analyses

Data for gross and histomorphological characteristics were analysed by analyses of variance for a completely randomized design using the General Linear Models (GLM) procedure of SAS (1987). The time of orchidectomy or location within the testis or their interactions with the type of line did not influence (P > 0.10) any histomorphological characteristic; thus the model included only line. The same analyses were used for oLH and hCG receptor characteristics. The main effect means were compared orthogonally (control line versus selected lines, and low line versus high line). Four rams did not exhibit an increase in testosterone secretion 2, 3 or 4 h after adding gonadotrophin compared with that after 1 h of incubation. These rams were considered non-responders and were not included in testosterone analyses. Data for testosterone secretion per hour per gram of testicular parenchyma were analysed by a split-split-plot design for each gonadotrophin using the GLM procedure of SAS (1987). The error term to test the type of line was animal within line. The error term to test dosage and the interaction of dosage with line was animal within line by dosage. This characteristic was also analysed with a model that included gonadotrophins. The main effects of line and gonadotrophin were tested using the error term animal within line by gonadotrophin by dosage.

The total testosterone secretion per gram of testicular parenchyma was calculated by summing the testosterone concentrations at each hour of incubation. The total testosterone secretion for each gonadotrophin was analysed by analysis of variance for a split-plot design. The main plot was line and the subplot included dosage and the interaction of line with dosage. Line was tested with animal within line as the error term. This characteristic was also analysed with a model that included gonadotrophin. The main effects were tested with animal within line by gonadotrophin as the error term.

Results

Gross and histomorphological characteristics

The scrotal circumference did not differ (P > 0.05) between trials or among lines and averaged 25.7 ± 1.6 (SD) cm. In Trial 1, paired testis mass, and in Trial 2, left testis mass did not differ (P < 0.05) among lines and averaged 189.3 ± 11.4 g and 91.5 ± 4.5 g, respectively. Values for right and left testicular tone did not differ (P > 0.05) among lines in Trial 2 and were 1.6 ± 0.18 and 1.6 ± 0.14 units, respectively.

The seminiferous tubule diameter did not differ (P > 0.05) among lines. Mean diameters and the number of seminiferous tubules counted were: 169.6 ± 7.5 μm, 158 ± 76; 178.2 ± 8.7 μm, 160 ± 31; and 182.2 ± 8.0 μm, 149 ± 26 for LL, HL and CL rams, respectively.

The testis volume and percentage volumes of seminiferous tubules, interstitium or components of the interstitium other than Leydig cells did not differ (P > 0.05) among lines (Table 1). The percentage volume of Leydig cells did not differ (P > 0.05) among lines; however, the percentage volume of vascular tissue was greater in LL (P < 0.05) rams than in HL or CL rams (Table 1). Percentage volumes of vascular tissue did not differ (P > 0.05) between HL and CL rams.

Leydig cell volume, the volume of Leydig cell nuclei, the number of Leydig cells per gram of testis, the total number of Leydig cells per testis, and the total volume of Leydig cells in the testis did not differ (P > 0.05) among lines (Table 2).

Receptor binding capacity and K_d values

The binding capacities of both [125I]-labelled oLH and [125I]-labelled hCG to testicular membrane preparations was saturable for rams in each line. Of the 23 rams evaluated, membrane preparations from only three rams exhibited saturation curves that could not be fitted directly to the binding equation for a one ligand, one binding site model.

The binding capacities of testicular preparations for oLH and hCG expressed as fmol mg^-1 of membrane protein, fmol g^-1 of parenchyma, or pmol per testis did not differ (P > 0.05) among lines (Table 3). The number of binding sites per Leydig cell for oLH and hCG did not differ (P > 0.05) among lines (Table 3). Capacity, regardless of how it was expressed, was greater (P < 0.05) for oLH binding than for hCG binding within each line of rams (Table 3). The capacity per gram of testicular parenchyma and the number of receptor sites per Leydig cell for oLH were approximately 1.5 times higher than for hCG.

Dissociation constants (K_d x 10^12 mol^-1 l^-1) for both oLH and hCG binding sites did not differ (P > 0.05) among lines (Table 4). The K_d values for oLH binding sites were approximately six times higher (P < 0.05) than for hCG binding sites within each line (Table 4).
Testicular traits and selection for reproductive rate

Table 2. Least-squares means for Leydig cell (LC) characteristics in Rambouillet rams from lines selected for low (LL) or high (HL) reproductive rates and from a random-bred control line (CL)

<table>
<thead>
<tr>
<th>Selection line</th>
<th>n</th>
<th>LC volume (µm³)</th>
<th>LC nuclei volume (µm³)</th>
<th>Total volume of LC (cm³)</th>
<th>Number of LC per testis (× 10⁶)</th>
<th>Number of LC per g of testis (× 10⁶)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LL</td>
<td>8</td>
<td>943.5</td>
<td>207.5</td>
<td>2.0</td>
<td>2.129</td>
<td>23.240</td>
</tr>
<tr>
<td>HL</td>
<td>6</td>
<td>1083.6</td>
<td>226.5</td>
<td>2.3</td>
<td>2.210</td>
<td>24.887</td>
</tr>
<tr>
<td>CL</td>
<td>6</td>
<td>917.5</td>
<td>203.9</td>
<td>1.9</td>
<td>2.015</td>
<td>23.813</td>
</tr>
</tbody>
</table>

SEM<sup>a</sup>

Table 3. Least-squares means for characteristics of LH and hCG testicular receptors of Rambouillet rams from lines selected for low (LL) or high (HL) reproductive rates and from a random-bred control line (CL)

<table>
<thead>
<tr>
<th>Selection line</th>
<th>n</th>
<th>Capacity per mg of membrane protein (fmol)</th>
<th>Capacity per g of parenchyma (fmol)</th>
<th>Capacity per testis (pmol)</th>
<th>Receptors per Leydig cell (× 10⁶)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>oLH&lt;sup&gt;a&lt;/sup&gt;</td>
<td>hCG</td>
<td>oLH&lt;sup&gt;a&lt;/sup&gt;</td>
<td>hCG</td>
</tr>
<tr>
<td>LL</td>
<td>6</td>
<td>45.3</td>
<td>17.2</td>
<td>383.2</td>
<td>147.1</td>
</tr>
<tr>
<td>HL</td>
<td>7</td>
<td>62.6</td>
<td>23.2</td>
<td>554.4</td>
<td>208.7</td>
</tr>
<tr>
<td>CL</td>
<td>7</td>
<td>48.1</td>
<td>19.9</td>
<td>378.8</td>
<td>157.6</td>
</tr>
</tbody>
</table>

SEM<sup>b</sup>

<sup>a</sup>Values are significantly different (P < 0.05) between oLH and hCG, within selection lines for each trait.

<sup>b</sup>Pooled SEM.

Table 4. Least-squares means for dissociation constants (K<sub>d</sub>) values for oLH and hCG testicular receptors of Rambouillet rams from lines selected for low (LL) or high (HL) reproductive rates and from a random-bred control line (CL)

<table>
<thead>
<tr>
<th>Selection line</th>
<th>n</th>
<th>K&lt;sub&gt;d&lt;/sub&gt; × 10&lt;sup&gt;-12&lt;/sup&gt; mol l&lt;sup&gt;-1&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>oLH&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>LL</td>
<td>6</td>
<td>87.3</td>
</tr>
<tr>
<td>HL</td>
<td>7</td>
<td>82.5</td>
</tr>
<tr>
<td>CL</td>
<td>7</td>
<td>133.8</td>
</tr>
</tbody>
</table>

SEM<sup>b</sup>

<sup>a</sup>Values are significantly different (P < 0.05) between oLH and hCG within each line.

<sup>b</sup>Pooled SEM.

Gonadotrophin-stimulated testosterone secretion in vitro

Testosterone secretion stimulated by oLH in vitro per gram of testis per hour did not differ (P > 0.05) among lines over time (Fig. 1a). However, there was a line by time interaction (P < 0.05) for hCG-stimulated testosterone secretion in vitro per gram of testis (Fig. 1b). LL and HL rams secreted more (P < 0.05) testosterone after 2 h of incubation compared with after 1 h. Concentrations of testosterone after 3 h did not differ (P > 0.05) relative to that after 2 h but had decreased at 4 h of incubation in LL and HL rams. Testosterone secretion in CL rams did not change (P > 0.05) during the first 2 h, but had decreased (P < 0.05) by 3 h and again (P < 0.05) by 4 h of incubation. During every hour of incubation HL rams secreted significantly more (P < 0.05) testosterone than did LL rams, with the exception of the value recorded after 3 h.

Testosterone secretion over time for equimolar dosages of oLH and hCG did not differ (P > 0.05) between gonadotrophins and there was no line by dosage by gonadotrophin interaction (P > 0.05) for testosterone secretion per hour.

Total oLH-stimulated and hCG-stimulated testosterone increased (P < 0.05) as the dosage of gonadotrophin increased (Table 5). The total amount of testosterone secreted per gram of testis did not differ (P > 0.05) between gonadotrophins at dosages of 0, 460 and 4600 pmol l<sup>-1</sup> (Table 5). However, 460 pmol hCG l<sup>-1</sup> stimulated the secretion of significantly more (P < 0.05) testosterone than did 0 pmol hCG l<sup>-1</sup>, whereas the same concentration of oLH did not significantly increase (P > 0.05) testosterone secretion compared with 0 pmol oLH l<sup>-1</sup> (Table 5). The type of line did not affect (P > 0.05) total testosterone secretion and there was no (P > 0.05) line by gonadotrophin interaction.

Discussion

Curry et al. (1993) reported that neither right, left nor paired testis mass or scrotal circumference differed among rams of
vascular tissue in LL rams influences testosterone secretion in vitro in LL rams was not evaluated in the present study.

Results of the binding studies of this experiment indicate that selection for or against reproductive rate in females of these lines has not altered oLH or hCG binding capacity, the number of Leydig cell receptor sites in, or affinity of receptor sites for, homologous or heterologous gonadotrophins (oLH or hCG) in males of these lines. This result is similar to that of Hochereau-de Reviers et al. (1990b), who reported that Ile de France rams (a prolific breed) and Romanov rams (a less prolific breed) do not differ in their LH-binding capacity per Leydig cell.

Within each line, the $K_d$ value for the binding of oLH to its receptors was approximately six times higher than the $K_d$ value for hCG, and binding capacities were approximately three times higher for oLH than for hCG. These results agree in part with those of Sairam et al. (1988), who found higher binding capacities for oLH than for hCG but similar $K_d$ values for both gonadotrophins in testicular preparations from rams. In contrast, Huhtaniemi and Catt (1981) reported similar binding capacities for oLH and hCG but a lower $K_d$ value for hCG than for oLH in rat testicular preparations. They observed that the binding of oLH in the rat fit a model with two different binding sites: one with a high affinity (low $K_d$) that constituted about 20% of the oLH receptors, and the other with low affinity (high $K_d$) that constituted 80% of the oLH receptors. In our study, binding data for hCG fit exclusively a one ligand, one-binding site model and, for the most part, so did the binding data for oLH, with the exception of three rams that appeared to fit a one ligand, two-binding site model.

The line by time interactions for hCG-stimulated testosterone secretion in vitro indicates that selection has altered the manner in which Leydig cells of rams of these lines respond to gonadotrophin stimulation over time. Although not significant at the $P < 0.05$ level, the same trend was apparent for oLH-stimulated testosterone secretion in vitro. HL rams secreted more hCG-stimulated testosterone during every hour of culture (except the third) than did LL rams. The mechanisms by which selection for reproductive rate in females has changed the pattern of response of the testis to hCG and possibly to oLH in vitro are not clear. It is known that testosterone

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### Table 5. Least-squares means for total testosterone secretion per gram of testicular parenchyma in vitro after stimulation by equimolar dosages of LH or hCG in Rambouillet rams

<table>
<thead>
<tr>
<th>Dosage of gonadotrophin (pmol l$^{-1}$)</th>
<th>Total oLH-stimulated testosterone secretion (ng g$^{-1}$ parenchyma)</th>
<th>Total hCG-stimulated testosterone secretion (ng g$^{-1}$ parenchyma)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>($n = 21$)</td>
<td>($n = 21$)</td>
</tr>
<tr>
<td>0</td>
<td>16.5$^a$</td>
<td>16.6$^a$</td>
</tr>
<tr>
<td>4.6</td>
<td>19.5$^a$</td>
<td>22.5$^b$</td>
</tr>
<tr>
<td>46</td>
<td>28.1$^c$</td>
<td>28.3$^c$</td>
</tr>
<tr>
<td>4600</td>
<td>35.6$^d$</td>
<td>34.9$^d$</td>
</tr>
<tr>
<td>SEM$^a$</td>
<td>5.7</td>
<td>5.7</td>
</tr>
</tbody>
</table>

$^a$Values within columns with different superscripts are significantly different ($P < 0.05$).

$^a$Pooled SEM.
Testicular traits and selection for reproductive rate

synthesis and secretion by the testicular parenchyma can be influenced by numerous intracellular and extracellular factors (Hall, 1988). Ewing et al. (1979) suggested that differences between certain mammals in LH-stimulated testosterone secretion are not related to differences in Leydig cell mass. Furthermore, Zirkin et al. (1980) found that differences in testosterone secretion among various species of male could be accounted for by differences in the volume of smooth endoplasmic reticulum in Leydig cells. The possibility that HL rams secrete more testosterone in vitro in response to hCG because their Leydig cells contain a greater volume of this type of endoplasmic reticulum than do Leydig cells of LL or CL rams remains to be examined.

Although the capacity of the oLH receptor was higher than the capacity of the hCG receptor, total oLH- and hCG-stimulated testosterone secretion did not differ in two out of three equimolar doses of gonadotrophin. It is possible that the higher affinity of the hCG receptor could have compensated for its lower capacity, thereby stimulating testosterone secretion to the same degree as oLH. Stimulation of testosterone secretion exhibited after administering the lowest dosage of hCG tested may be explained, at least partially, by the higher affinity of hCG for its receptor, which, according to Huhtaniemi and Catt (1981), contributes to the well-known high bioactivity of this hormone. These results also indicate that the ram testis seems to be more sensitive to hCG than to oLH.

In conclusion, selection for or against reproductive rate in Rambouillet ewes has not altered gross or most histomorphometric characteristics of the testis of male offspring, with the exception that selection against reproductive rate increases the proportion of testicular volume occupied by vascular tissue within the interstitium. Furthermore, selection has not altered total oLH- and hCG-stimulated testosterone secretion in vitro or testicular oLH and hCG receptor affinity or capacity of male offspring.

This paper is contribution no. J-2816 of the Montana Agricultural Experiment Station. The research was supported in part by an SR-CSRP grant from the Agency for International Development, Title XII, Grant no. AID/DSANXII-G-0049. Ovine LH and hCG were generously provided by the NIDDK, the National Hormone and Pituitary Program, and the University of Maryland Medical School.

References

Abercrombie M (1946) Estimation of nuclear population from microtome section Anatomical Record 94 239-249

Byeley DJ, Bertrand JK, Berardinelli JG and Kiser TE (1990) Testosterone and luteinizing hormone response to GnRH in yearling bulls of different libido Theriogenology 34 1041–1049
Carr WR and Land RB (1975) Plasma luteinizing hormone levels and testis diameter of ram lambs of different fertility Journal of Reproduction and Fertility 42 325–333
Curry KC, Berardinelli JG, Burfening PJ and Adair R (1993) Selection for reproductive rate in rams and feeding regimen on testicular traits and epididymal sperm reserves in Rambouillet rams Small Ruminant Research 11 257–265
Fawcett DW, Neaves WB and Flores M (1973) Comparative observations on interstitial lymphatics and the organization of the intratesticular tissue of the mammalian testis Biology of Reproduction 9 500–532
Hoffmann EO, Flores TR, Couver J and Garret HB (1983) Polychrome stain for high resolution light microscopy Laboratory Medicine 14 779–781
Schoenian CS and Burfening PJ (1990) Ovulation rate, lambing rate, litter size and embryonic survival in Rambouillet sheep selected for high and low reproductive rate Journal of Animal Science 68 2263–2270

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