Body weight, semen quality, and age at sperm production of hema-castrated and intact cockerels (Gallus domesticus)

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Summary. Spermatozoa were found earlier in the ejaculates of hema-castrated cockerels and the mean body weight of the hema-castrates was significantly (P < 0.01) greater than that of the intact cockerels. The mean values for sperm concentration/ml, and total spermatozoa in the ejaculates of the hema-castrates were significantly greater (P < 0.01) than those of the intact cockerels. Differences existed between the hema-castrates bearing the left testis, and those bearing the right testis in respect of age at sperm production, sperm/concentration and total spermatozoa in the ejaculate.

Introduction

Hematicastration increases plasma gonadotrophin concentration in cockerels (Wilson & de Reviers, 1979), with consequent enhancement of spermatogenesis (Onuora, 1985). Differences in size of the left and right testis have been observed in cockerels (Witschi, 1935; Romonoff, 1960). Witschi (1935) suggested that this difference was due to the larger number of germ cells which migrate to the primordial left testis on the 4th day of incubation. Onuora (1987) determined when spermatozoa were first found in the macerated tissues of the testes of hema-castrates bearing the left testes, hema-castrates bearing the right testes and intact cockerels, and reported that spermatozoa were found earlier in these tissues in the following order: left testis remaining in hema-castrate, right testis remaining in hema-castrate, left testis of intact cockerel and right testis of intact cockerel. Removal of the right testis might then be expected to lead to early sperm production.

The aim of the present study was to determine when spermatozoa could be obtained from the ejaculates of hema-castrated and intact cockerels, and to compare the quality of the semen obtained from these cockerels.

Materials and Methods

Commercial Babcock chicks were reared in deep litter and fed broiler starter (containing 21.0% crude protein, 3.2% oil, 4.2% fibre and supplemented with vitamins and minerals) and water freely. At 5 weeks of age just before hema-castration the birds were arbitrarily divided into three groups of 15 each. Hematicastration was carried out surgically after infiltration of 2% xylocaine (Astra, Sweden) into the last intercostal space. All the birds were hema-castrated within 24 h, alternating between birds in Group HLT and birds in Group HRT. During hematicastration there were some deaths. Finally each group was made up of 10 birds having a mean body weight of 0.41 ± 0.05 for hema-castrates bearing the left testis (Group HLT), 0.39 ± 0.15 for hema-castrates bearing the right testis (Group HRT) and 0.42 ± 0.08 kg for intact birds (Group CT). At 10 weeks of age all the cockerels were put into individual cages and fed broiler finisher diet (containing 19.0% crude protein, 4.0% oil, 4.2% fibre, and supplemented with vitamins and minerals) and water freely until the end of the study.

From 12 weeks of age ejaculates were collected at intervals of 2 days (i.e. 3 times a week) using the massage technique of Burrows & Quinn (1935). The mean body weight of the groups was determined weekly from 12 weeks of age. Initially smears made from the ejaculates were stained with eosin/nigrosin, and searched for the presence of spermatozoa (Cooper & Rowell, 1958). When spermatozoa had been found in all the ejaculates of all the cockerels, the semen samples were thereafter evaluated for sperm concentration, motility, percentage of abnormal spermatozoa, total spermatozoa in the ejaculate and volume of semen. In total, 18 ejaculates were collected before and 33 after spermatozoa were found in the ejaculates of all the birds. The latter data were analysed using Student’s t test.
Table 1. Number of hemicastrated and intact cockerels and age at which spermatozoa were found in their ejaculates

<table>
<thead>
<tr>
<th>Age (days)</th>
<th>Group HRT (N = 10)</th>
<th>Group HLT (N = 10)</th>
<th>Group CT (N = 10)</th>
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<tr>
<td>111</td>
<td>7</td>
<td>3</td>
<td>0</td>
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<td>121</td>
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<td>6</td>
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<tr>
<td>147</td>
<td>10</td>
<td>—</td>
<td>8</td>
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<tr>
<td>150</td>
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<td>—</td>
<td>10</td>
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</table>

Ejaculates were collected 3 times a week from hemicastrates bearing the right testis (Group HRT), hemicastrates bearing the left testis (Group HLT) and intact cockerels (Group CT).

Fig. 1. The body weights of hemicastrated and intact cockerels at different ages. Each value represents the mean ± s.e.m. of 10 observations.

Results

The ages at which spermatozoa were found in hemicastrated and intact cockerels are shown in Table 1. Spermatozoa were first observed in the ejaculates of the hemicastrates at 111 days and in the intact cockerels at 121 days of age.
Figure 2. Values of (a) semen volume, (b) percentage abnormal spermatozoa, (c) percentage motility of spermatozoa, (d) total numbers of spermatozoa and (e) sperm concentration in hemicastrated and intact cockerels at different ages. Values are the mean of 10 observations.

Figure 1 shows the mean of the body weight of hemicastrated and intact cockerels at different ages: between 12 and 28 weeks of age values were higher in hemicastrates ($P < 0.01$) than in the controls. There was no significant difference between the body weights of birds in Groups HRT and HLT.

There was no significant difference between the groups in respect of the volumes of the ejaculates (HRT $0.28 \pm 0.06$; HLT $0.22 \pm 0.01$; CT $0.22 \pm 0.03$ ml) (Fig. 2a), and percentage of
abnormal spermatozoa (HRT 1.67 ± 0.2; HLT 1.17 ± 0.25; CT 1.62 ± 0.16) (Fig. 2b), and percentage motility (HRT 77.8 ± 0.35; HLT 77.4 ± 0.37; CT 78.28 ± 0.50) (Fig. 2c).

Sperm concentration (Fig. 2e) increased with age. There was a significant difference between the hemicastrated and the intact cockerels (P < 0.01) and between Groups HLT and HRT (P < 0.01) (HRT 1.61 ± 0.11; HLT 1.27 ± 0.26; CT 1.16 ± 0.26 spermatozoa/ml × 10^9).

Figure 2(d) shows that the mean total spermatozoa in the ejaculate of each group increased with age. There was a significant difference between the hemicastrates (HRT 0.449 ± 0.086 × 10^9; HLT 0.350 ± 0.064 × 10^9) and the controls (0.287 ± 0.054 × 10^9) (P < 0.01), and between the birds in Groups HLT and HRT (P < 0.01).

Discussion

The results of this study show that hemicastrated cockerels produced spermatozoa earlier than did intact cockerels and that all the birds in Group HLT produced spermatozoa before those in Group HRT. These results confirm an earlier report (Onuora, 1987) in which spermatozoa were found in the macerated tissues of the testes of all Group HLT birds earlier than in those of Group HRT, and in those of Group HRT earlier than those of the intact cockerels. The mechanism for the earlier production of spermatozoa is not understood but may be associated with the enhanced spermatogenesis in hemicastrates (Santolaya & Burgos, 1978; Onuora, 1985) and to differences in the attributable number of 'spare' (functional) receptors and occupancy of LH receptors in the Leydig and Sertoli cells (Sharpe, 1982).

The difference between Groups HRT and HLT may be due to anatomical features. In chicken the number of blood vessels to the left and right gonad is related to their activity (Nickel et al., 1977). Furthermore, at the 4th day of incubation 4-5-fold the number of germ cells migrate to the primordial left testis than to the right testis (Witschi, 1935). The differences may be due to the spatial disposition of the testes with respect to the air sacs, as there have been suggestions that the testes are cooled by these air sacs during thermal control of spermatogenesis (King & McLelland, 1975).

The mean body weights of the hemicastrates were significantly greater than those of the controls, and Group HRT birds were slightly heavier than those in Group HLT, an observation already reported (Mast et al., 1981; Onuora, 1987). However, Wilson & de Reviers (1979) did not find any significant difference between the body weights of hemicastrated and intact cockerels.

Hemicastration causes relaxation of the pituitary–gonadal and Sertoli cell–inhibin axes, causing initial reduction in the amount of inhibitor and testosterone plasma concentrations (Walton et al., 1978; Droit et al., 1979; de Krester, 1979). The resultant higher concentrations of LH, FSH and testosterone (Franchimont et al., 1978; Cunningham et al., 1978; Wilson & de Reviers, 1979; Walton et al., 1980) increase the production of both intra- and extra-tubular components and their precursors in the prepubertal testis (de Jong & Sharpe, 1977; Cunningham et al., 1978). The increased spermatogenic components enhance spermatogenesis with production of more spermatozoa both within the testis (Thyagaraja & Sakar, 1970; Santolaya & Burgos, 1978; Hochereau-de Reviers & Courrot, 1978; Onuora, 1987) and extragonadally (Onuora, 1985). Hence the sperm concentration, and the total spermatozoa in the ejaculate, were significantly higher in the hemicastrates than in the intact birds. The rise in the androgen concentration may be responsible for the increased body weight in the hemicastrates.

The difference in the degree of testicular compensatory hypertrophy and the concomittant plasma concentrations of androgen between the hemicastrated and the intact cockerels may explain the presence (Mast et al., 1981; Putra & Blackshaw, 1982; Onuora, 1987; present study) or absence (Wilson & de Reviers, 1979) of significant differences between the body weights. However, plasma testosterone concentrations correlate positively with body weights (Gilmore, 1969; Gemmell et al., 1986).
The results from this study show that the left and right testes have different spermatogenic potentials. Although the left testis produced spermatozoa earlier, the right testis has more spermatogenic activity when total spermatozoa per mg testis are compared. The mechanisms responsible for these differences are not known and need further investigation.

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References


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