Testis development and daily sperm output in guinea-fowl raised under constant daily photoperiods

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Summary. The testicular growth of guinea-fowls in relation to age was compared under 3 constant photoperiods (7L : 17D ; 14L : 10D ; 20L : 4D) from 3 weeks of age. Although long daylength (14L or 20L) caused precocious development of the gonads, it also limited total adult gonadal weight to around 1 800 ± 200 mg. Short daylengths (7L) delayed the development of the gonads but their adult weight was enhanced to 2 800 ± 200 mg.

A comparison of the changes in daily sperm output (DSO) under 7L : 17D and 14L : 10D was consistent with the observations on testis weights. Under neither photoschedule was a correlation found between individual age at first ejaculate and the individual DSO observed at sexual maturity. However, the mean individual DSO’s estimated from the first 10 ejaculates were correlated to those observed at sexual maturity, and the latter were also correlated to the DSO’s observed at 53-55 or 59-61 weeks of age.

Introduction.

Reports on reproduction in the guinea-fowl are not numerous although it has been domesticated for a long time and is an important agricultural product. Barbier et Leroy (1970) described seasonal variations in the testicular development of guinea-fowls fed ad libitum but submitted to natural climatic conditions (geographical location of the experiment : latitude 49° N). Maximal gonadal development was observed during the summer (May to September : 2.5 ± 1.0 g) and the minimum appeared between October and January (0.5 ± 0.5 g).

In equatorial regions of Africa, reproduction in wild species of guinea-fowls occurs at any time during the year, but the appearance of a breeding season is more marked as the latitude increases (for example, it only occurs between March and May in Morocco : Barbier and Leroy, 1970).

These observations support the idea that photoperiodism may be an important factor regulating reproduction in the guinea-fowl. The present paper describes for the first time changes in testis weight and sperm output under short and long constant photoperiods in controlled environmental conditions.

Materials and methods.

Experiment 1. — 360 one-day-old male guinea-fowls were divided into 3 batches (120 chicks in each). All the animals were fed and allowed water ad libitum (table 1)
throughout the experiment which lasted 70 weeks. They were reared on the ground till the age of 16 weeks, then individually caged. The ambient temperature was maintained at 35 °C until 3 weeks of age, at 32 °C up to 6 weeks and at 20 °C thereafter. The photoperiod was gradually reduced to either 20L : 4D (at 1 week of age, group 1), 14L : 10D (at 2 weeks of age, group 2) or 7L : 17D (at 3 weeks of age, group 3). These daily photoperiods were then respectively maintained until the end of the experiment (throughout the text, L refers to the hours of illumination and D to the hours of darkness).

The animals were individually weighed at various ages, from the 8th to the 51st week after hatching. At 8, 12, 16, 20, 24, 28, 34, 39, 51 and 70 weeks of age, 10 animals in each group were killed and autopsied. The testes were immediately weighed to the nearest mg.

Experiment 2. — Using similar conditions of temperature, feeding and housing, 2 of the above photoschedules (14L : 10D and 7L : 17D) were replicated and 72 males were submitted to each. In these animals, semen collections were performed twice a week at various periods between 22 and 61 weeks of age by a massage technique derived from Burrows and Quinn (1935).

This frequency of semen collection allows the maximum number of ejaculated spermatozoa per male to be obtained (table 2).

**TABLE 1**

*Composition of the diets distributed « ad libitum » in the 3 trials*

<table>
<thead>
<tr>
<th></th>
<th>0-6 weeks (starting period)</th>
<th>7-16 weeks (growing period)</th>
<th>17-61 weeks (adult period)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Metabolizable (kcal/kg)</td>
<td>3 150</td>
<td>3 000</td>
<td>2 600</td>
</tr>
<tr>
<td>Energy (MJ/kg)</td>
<td>13.2</td>
<td>12.5</td>
<td>10.9</td>
</tr>
<tr>
<td>Crude protein Level</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(N x 6.25)</td>
<td>24</td>
<td>18</td>
<td>14</td>
</tr>
</tbody>
</table>

**TABLE 2**

*Variations in the sperm output as a function of the semen collection frequency*

<table>
<thead>
<tr>
<th>Frequency of collection (times a week)</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of males * collected</td>
<td>54</td>
<td>54</td>
<td>54</td>
</tr>
<tr>
<td>Number of spermatozoa ejaculated each week (in millions) **</td>
<td>174</td>
<td>371</td>
<td>361</td>
</tr>
</tbody>
</table>

* All males, raised under the same environmental conditions were collected over four consecutive weeks. The number of sperm is expressed as the mean of the 3 last weeks.
** F = 20 for 2, and 160 DF, highly significant differences appear between 1 vs 2 and 3 semen collections per week. No significant differences were observed between 2 and 3 semen collections/week.

Each ejaculate was individually weighed to the nearest mg and the results converted into volume (specific gravity of semen ~ 1 mg/μl). This method was found to be
more precise and quicker than pipetting because of the small volume of the ejaculates produced by guinea-fowl (range: 0-200 μl).

The sperm concentration of semen was evaluated by optical density at 600 nm (fig. 1). This allowed the total number of spermatozoa per ejaculate to be calculated, from which the daily sperm output (DSO) (Amann, 1970) was obtained. A highly significant correlation was observed between DSO and testicular weight (fig. 2).

**FIG. 1.** — Relationship between the total number of spermatozoa (counted with hemocytometers *) and optical density measured in the same ejaculates. * 10 replications per ejaculate. Total number of ejaculates = 60. Wavelength: 600 nm. Equation of regression: \( y = 0.170 + 0.0056x \) (\( r = 0.96 \)).

**FIG. 2.** — Relationship observed between the total testis weight and the number of ejaculated spermatozoa in guinea-fowl. All the males, placed under a 14L/10D light regime, were collected twice a week between 66 and 68 weeks of age. — Linear regression: \( Y = 1753.5 + 2748.5X \) (\( r = 0.84 \)).
As the sperm reserves contained in the deferent ducts may introduce bias when estimating sperm production by sperm output (Ortavant, 1958; de Reviers, 1972), the males were ejaculated twice during the week preceding each period of ejaculate measurement.

Experiment 3. — The mean individual DSO was estimated for 3 different periods in 15 and 45 males raised under 7L : 17D and 14L : 10D, respectively. Period I involved the first 10 ejaculates from individuals (mean respective ages at start: 27-28 and 21-22 weeks). Period II involved consecutive ejaculates from individuals at the end of the testicular growth phase (corresponding ages at start: 46-47 and 24-26 weeks). Period III involved 6 similar ejaculates between 53 and 61 weeks of age.

Results.

Experiment 1.

1. Body weight was quite similar under each photoschedule until 28 weeks of age; it decreased unexpectedly between 20 (1 600 ± 50 g) and 24 weeks (1 480 ± 50 g) of age under each photoschedule, and then increased again, the highest levels occurring

![Graph](image-url)
Photoperiodism and reproduction in guinea-fowl

under 7L : 17D and the lowest under 20L : 4D (1 900 ± 50 g vs 1 600 ± 30 g at 51 weeks of age). Under 14L : 10D and 20L : 4D, body weight appeared to be stabilized at 39 weeks of age.

2. Testicular development (fig. 3) was very similar under 20L : 4D and 14L : 10D. After a rapid increase from 8 to 20 weeks of age (60-80 mg up to 1 400-1 600 mg), the mean testicular weight was stable up to 51 weeks of age. It decreased thereafter only under the 20L : 4D photoschedule (1 240 ± 100 mg at 70 weeks of age).

In contrast, the testes developed much later under 7L : 17D, reaching their maximum weight at 51 weeks of age. This weight was higher than the maximum observed under long photoperiods (2 800 mg vs 1 800 mg), but it was not statistically different from that observed at 34, 39 and 70 weeks under the same treatment (F = 0.99).

Experiment 2. — The number of spermatozoa collected, expressed on a per day basis (daily sperm output), is shown in figure 4 (for 14L : 10D and 7L : 17D photoschedules). The DSO reached 30 × 10⁶ spermatozoa at 23 weeks of age in males maintained under 14L : 10D compared to 28 weeks of age in males maintained

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**FIG. 4.** — *A comparison of the changes in daily sperm output (DSO) of guinea fowls placed under 7L/17D or 14L/10D photoperiods. Each point represents the mean of two consecutive weeks ± SEM. Photoschedules: •—• 7L/17D; •—• 14L/10D.*

Reproduction, nutrition, développement
under 7L : 17D. Afterwards, under 14L : 10D, there was a slow, linear increase in the DSO from 23 to 55 weeks of age (30 to $64 \times 10^6$ sperm). In animals exposed to 7L : 17D, the DSO reached a maximum of $84 \times 10^6$ sperm/day at 38 weeks of age, and then slowly decreased until the end of the experiment.

**Experiment 3.** — Table 3 shows the results of the DSO observed in the same animals during 3 different age periods. A significant correlation ($P<0.05$) was obtained between periods I and II and III in both groups and between periods I and III in the 14L : 10D group. On the other hand, the ages at which the first ejaculate for each male were obtained were not found to be either positively or negatively correlated with adult or late individual sperm production in either group ($r < 0.25$).

**Table 3**

<table>
<thead>
<tr>
<th>Stage of development</th>
<th>Correlation between stages (individual DSO)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>I</td>
</tr>
<tr>
<td>Early sperm production</td>
<td>10</td>
</tr>
<tr>
<td>Sexual maturity</td>
<td>27-28</td>
</tr>
<tr>
<td>Later adult</td>
<td>23 ± 4</td>
</tr>
</tbody>
</table>

7L : 17D (15 males)

- No. of ejaculates/male: 45
- Mean age (weeks) at start: 21-22
- Mean DSO ($\times 10^6$/day): 37 ± 3

14L : 10D (45 males)

- No. of ejaculates/male: 45
- Mean age (weeks) at start: 21-22
- Mean DSO ($\times 10^6$/day): 37 ± 3

**Discussion.**

It may be concluded from the above results that testicular development and sperm output in the guinea-fowl are strongly influenced by daylength when constant daily photoperiods are used. These photoperiods, however, do not induce considerable modifications in body growth, as in the cockerel (Parker and Mc Cluskey, 1964; de Reviers, 1974) and the turkey (Krueger et al., 1977).

The variations in the testicular development observed under either short or long constant daylengths in the guinea-fowl are more similar to those reported in the turkey (Krueger et al., 1977) than in the cockerel. In the latter species, using Rhode × Wyandotte M519 cockerels, the rapid phase of testicular growth started at the same age (16 weeks) under 16L : 8D or 8L : 16D. Furthermore, the difference in the ages at which testicular growth terminated under these photoperiods was only 2-4 weeks (de...
In turkeys raised under 8L : 16D, the mean testicular weight observed at 30 weeks of age was only half of that observed at the same age under 15L : 9D (14.2 ± 7.9 g vs 33.5 ± 13.4, Krueger et al., 1977). This implies a difference between species either in the age at which the rapid phase of testicular growth starts, or in the growth rate of the testes, or both. At the end of testicular growth, the difference in the mean testicular weights observed in cockerels raised under 8L : 16D or 16L : 8D was clearly noticeable (16.8 ± 1.4 vs 19.5 ± 0.8 g at 24 weeks of age, de Reviers et al., 1974). A positive effect of long constant photoperiods on the level of testicular development in the adult cockerel was observed. In contrast, the adult testis weight of turkeys raised under long days (15L : 9D) was much lower than under short days (8L : 16D) (36.5 ± 6.4 vs 66.2 ± 20.2 g ; Krueger et al., 1977).

There appears to be, therefore, a strong species × daylength interaction with respect to testicular development under constant daily photoperiods. This might result from divergencies in photosensitive thresholds, which are known to be very different between wild and domestic species. The white-crowned sparrow, for instance, does not show testicular growth when submitted to a daily photoperiod of less than 9 hrs of light per day (Farner, 1957), while in the cockerel, development to the adult condition can occur, though slowly, even when the daily photoperiod is restricted to half an hour (Nalbandov, 1970). However, it is also known that testicular growth results from a coincidence between the light and a photosensitive phase occurring during the 24 hr. cycle (Bunning, 1936 ; Follett and Sharp, 1969) and subjected to circadian periodicity. The time at which this phase occurs and its duration, might vary between different species of birds as it does in mammals (Ortavant, 1977).

Another problem is the apparent antagonism between the precocious development of the testis and its adult weight found here in the guinea-fowl and, elsewhere, in the cockerel (de Reviers, 1975) and the turkey (Krueger et al., 1977). This antagonism, occurring under various photoschedules (constant daylengths in the guinea-fowl and turkey, but increasing daylengths in the cockerel) is in fact observed between photoschedules. Within a given photoschedule, the results of this study of DSO in the guinea-fowl do not show any correlation between age at first ejaculate and the DSO level observed when testicular development is terminated. The situation is probably the same for testicular development as for DSO.

Furthermore, wide individual variations in testicular weight and DSO were observed in the guinea-fowl irrespective of age and photoperiod, a situation similar to that in the cockerel (de Reviers and Williams, 1981). In both species, the individual DSO results obtained during the growth of the testes were well correlated with those observed at sexual maturity ; this presents an interesting possibility for selecting males before they are used for breeding purposes. The ranking of the males might result, at least partly, from the sizes of Sertoli and stem spermatogonia populations which are determined during the prepuberal period.

Résumé. Le développement testiculaire de pintades a été comparé sous 3 photopériodes quotidiennes constantes (7L : 17N ; 14L : 10N ; 20L : 4N) appliquées à partir de 3 semaines d'âge. Alors que les jours longs (14L : 10N) provoquent un développement précoce des testi-
cules, ils ont aussi pour conséquence de diminuer leur poids total à l'âge adulte aux environs de 1 800 ± 200 mg. Les jours courts (7L) retardent le développement des testicules mais leur poids total à l'âge adulte atteint 2 800 ± 200 mg.

Les variations du nombre de spermatozoïdes récoltés sous 7L ou 14L sont en bon accord avec celles observées pour le poids des testicules. Sous ces deux photopériodes, l'âge au premier éjaculat et les nombres moyens de spermatozoïdes récoltés à l'âge adulte ne sont pas corrélés. Par contre, les nombres de spermatozoïdes contenus dans les 10 premiers éjaculats sont corrélés avec les nombres de spermatozoïdes récoltés vers la fin du développement des testicules et ces derniers sont eux-mêmes corrélés avec ceux observés à 53-55 (14L) ou 59-61 (7L) semaines d'âge.

References

FARNER D. S., 1957. Avian photoperiodic testicular response and function of the hypothalamo-hypophysial axis. The Physiologist, 1, 26 (abstr.).