OESTRADIOL CONCENTRATION IN THE PERIPHERAL PLASMA OF THE DOMESTIC HEN FROM 7 WEEKS OF AGE UNTIL THE TIME OF SEXUAL MATURITY

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Summary. A specific radioimmunoassay technique was used to measure the concentration of oestradiol at weekly intervals in the blood plasma of seven pullets from 7 weeks of age until they were sexually mature. Large differences in oestradiol concentration were observed between birds. The mean concentration was 94 pg/ml plasma 7 weeks before the first egg was laid. The concentration then began to rise sharply 2 to 3 weeks later to reach a peak of mean value 355 pg/ml 2 to 3 weeks before laying began. The mean concentration when egg-laying was established was 138 pg/ml. The high oestradiol levels found before the onset of laying may be essential for the synthesis of yolk protein precursors and for the conservation of calcium as medullary bone in preparation for egg-shell formation. The peak may also be an important part of the changes which occur at puberty in the interrelationship between the hypothalamus, the pituitary gland and the ovary.

INTRODUCTION

Considerable changes in the blood composition of immature chickens or adult cockerels occur after oestrogen administration (Lorenz, 1954; Urist, 1959). The appearance in the plasma of the egg-yolk protein precursors, phosvitin and lipovitellin (Gruber, 1972), and the increase in the total calcium concentration are of the most direct relevance to reproduction. In the presence of androgens, oestrogens cause calcium to be deposited and stored as medullary bone (Benoit & Clavert, 1945), a soft bone tissue which invades the marrow cavity of long bones (Kyes & Potter, 1934) and which acts as a buffer against large fluctuations in calcium requirements (see Simkiss, 1967). All of these components as well as others under oestrogenic control, such as plasma proteins and lipids, increase in concentration as hens become sexually mature (Vanstone, Maw & Common, 1955; McIndoe, 1959; Heald & Badman, 1963; Redshaw & Follett, 1972). The finding that oestrone reaches a peak level in urine a few days before egg-laying begins (Common, Ainsworth, Hertelendy & Mathur, 1965; Mathur, Anastassiadis & Common, 1966) is in agreement with
the view that these changes occur under the influence of endogenous oestrogens. The hormonal interrelationships between the ovary, the pituitary and the hypothalamus in the hen are still far from clear (Gilbert, 1971) and the changes in these interrelationships during sexual maturation are virtually unknown.

An examination of the blood oestradiol levels up to the time of sexual maturity was therefore undertaken to determine whether changes in oestradiol concentration accompany the metabolic changes which occur during maturation, and to provide some information on the hormonal regimen under which ovulation commences.

MATERIALS AND METHODS

Seven birds (Sussex White pullets × Rhode Island Red cockerels), hatched on 12th November 1971, were reared to sexual maturity. From 7 weeks of age, they were accommodated in three adjacent wire cages and were subjected to 14 hr light/day throughout the entire rearing period. They were fed a chick mash up to 11 weeks of age, after which they received a standard grower’s ration.

Samples of blood (2 ml) were removed from each bird between 09.30 and 10.30 hours at 7 and 10 weeks of age and weekly samples were removed thereafter until 1½ to 2 weeks after the first egg was laid. The successive samples were removed from alternate brachial veins into a heparinized syringe. The blood was immediately cooled to 4°C and the plasma was separated and stored at −20°C until it was assayed. All samples from a given bird were assayed at the same time to eliminate interassay variation.

Oestradiol was assayed by the radioimmunoassay procedure described previously (Senior, 1973, 1974). Plasma (0.5 to 1.0 ml), to which was added 16,000 d/min (20 pg) of [3H]oestradiol-17β, was extracted with three 5-ml quantities of diethyl ether. The dried ether extract was chromatographed on a 15-cm long column of Sephadex LH-20, using the solvent system 15% methanol and 85% toluene. The fraction of the eluate containing oestradiol was evaporated to dryness and incubated overnight at 4°C with a diluted antiserum preparation. An aliquot of the incubate was removed to estimate the total radioactivity in the assay tube and also to determine the loss of oestradiol throughout the procedure. Bound and free radioactivity were separated using dextran-coated charcoal. The values for the percentage of radioactivity bound to the antiserum were compared with those for standard quantities of oestradiol-17β. The assay was able to detect concentrations as low as 20 to 30 pg/ml plasma. In six of the birds, single estimates were made on 1 ml plasma. Samples from Bird M5 were assayed twice, using 0.5 ml plasma for each assay.

The antiserum was prepared in sheep by injecting oestradiol-17-succinyl bovine serum albumin. It was capable of binding oestrone, oestradiol-17α and oestriol to an extent which was 80%, 20% and 10%, respectively, of the binding of oestradiol-17β. The binding of all other steroids tested was less than 0.01% of that of oestradiol-17β. The Sephadex LH-20 column procedure separated oestrone and oestriol from oestradiol-17β and oestradiol-17α, but the latter two oestrogens were not separated from each other. The oestradiol results presented, therefore, may include a contribution by oestradiol-17α.
RESULTS

The age at which each bird laid the first egg is shown in Table 1. The first egg was never laid in isolation. The birds immediately began laying clutches of two to six eggs separated by inter-clutch intervals of only 1 day. Birds M1 and M5 began laying at 17½ weeks of age whereas the other five birds laid their first egg between 20 and 22 weeks of age.

Table 1. Concentrations of oestradiol in the peripheral plasma of hens from 7 weeks of age until sexual maturity

<table>
<thead>
<tr>
<th>Age (weeks)</th>
<th>Bird</th>
<th>M1 (122 days)</th>
<th>M2 (147 days)</th>
<th>M3 (149 days)</th>
<th>M4 (141 days)</th>
<th>M5 (122 days)</th>
<th>M6 (142 days)</th>
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The oestradiol concentrations are expressed in terms of pg/ml. The age at which the first egg was laid is shown in parentheses after each bird number. The arrow represents the approximate occurrence of the onset of laying.

The peripheral plasma concentrations of oestradiol are shown in Table 1. Text-figure 1 shows the combined results, taking the time at which the first egg was laid as the reference point. Oestradiol levels began to rise sharply 4 to 5 weeks before the first oviposition, reached a peak 2 to 3 weeks beforehand and fell rapidly in the 2 weeks before egg-laying began. The peak was clearly shown in all birds except M4. The significant nature of this peak in a single bird was emphasized by the duplicate results obtained for Bird M5.

The large differences in oestradiol concentration between birds are reflected in the large standard errors (Text-fig. 1). For example, the maximum concentration of oestradiol in Bird M7 was almost 1 ng/ml whereas that in Bird M4 was only 160 pg/ml. There was also a difference in the pattern of levels with age between birds, although they all showed the highest levels just before commencing to lay. Four birds (M1, M5, M6 and M7) showed a single distinct peak whereas the other three had subsidiary peaks or a prolonged peak. The mean peak level (355 pg/ml) was significantly higher than the level 7 weeks before the onset of laying (94 pg/ml, P<0.02) and also than the level found subsequently during the laying period (138 pg/ml, P<0.05). The mean level
TEXT-FIG. 1. Concentration of oestradiol in the peripheral plasma of pullets approaching sexual maturity. Data from seven birds were pooled with reference to the week during which the first egg was laid. The mean (± S.E.) for each time period is shown.

found after egg laying began was significantly higher than that before 10 weeks of age ($P<0.001$).

**DISCUSSION**

Pre-laying peaks of several blood constituents have been observed in maturing hens. The concentration in plasma of total proteins (Vanstone et al., 1955), total lipids and fatty acids (Heald & Badman, 1963) and phospholipoproteins (McIndoe, 1959; Heald & Badman, 1963) reaches a peak a few days before the onset of laying. These constituents probably, therefore, reach their maximum concentration some time after the oestradiol peak observed in this investigation. Redshaw & Follett (1972), however, found that the yolk-protein precursor, lipovitellin, whose synthesis by the liver is specifically oestrogen-dependent (Gruber, 1972), reached a peak concentration in plasma 1 to 2 weeks before the onset of laying. The timing of this peak is therefore close to that of oestradiol found in this study. The physiological significance of these observations in relation to the development of the ovarian follicles remains to be elucidated.

Oestrogens (in the presence of androgens) cause an increase in the retention of dietary calcium (Common, Rutledge & Hale, 1948) and induce medullary bone formation (Benoit & Clavert, 1945). In maturing birds, medullary bone develops 10 to 14 days before the first egg is laid and the retention of calcium increases from about 100 mg/day to as much as 750 mg/day before laying commences (see Taylor, 1965). The medullary bone is markedly depleted once
egg production begins (Cox & Balloun, 1971). The pre-laying peak of oestradiol may, therefore, help in building up the body reserves of calcium in preparation for egg-shell formation.

Common et al. (1965) and Mathur et al. (1966) found that the highest content of oestrone in urine occurred only ‘a few days’ before laying began. Thus, the time when urinary oestrone levels are high may coincide with the time when plasma oestradiol levels are falling. Such a time relationship, though requiring confirmation in a single experiment, points to the possibility of a marked change in either the metabolism or the excretion rate of oestrogens shortly before the onset of laying. In experiments involving the injection of radioactive oestradiol-17β and oestrone and the determination of urinary metabolites, Common, Mathur, Mulay & Henneberry (1969) noted that the equilibrium between oestradiol-17β and oestrone was considerably more in favour of oestrone in laying than in non-laying hens. Mathur & Common (1969) found relatively more oestrone than oestradiol-17β in urine from laying hens than from non-laying hens. This evidence suggests that oestrone becomes relatively more, and oestradiol-17β relatively less important as birds come into lay. Whether such a change is sufficient to account for a simultaneous fall in blood oestradiol levels and rise in urinary oestrone content remains to be determined.

Recent evidence (Cunningham, Bonney, Furr & Onuora, 1973; P. J. Sharp, personal communication) suggests that the concentration of LH in the peripheral plasma of pullets rises from 16 weeks of age onwards and may reach a maximum value before the onset of laying. Cunningham et al. (1973) also found that the quantity of LH released after an injection of synthetic LH-releasing factor (LH-RF) declined to zero between 17 and 21 weeks of age. Just before pullets come into lay, therefore, the rate of synthesis and release of LH may be maximal. The rôle of ovarian steroids in the control of LH release in the hen is still far from clear and the significance of these hormonal changes during the onset of sexual maturity must await further investigation.

The reason for the large variation between birds is not entirely clear although the infrequent sampling intervals may account for some of the differences. The stage at which the ovary begins its cyclic follicular development is uncertain. Wood-Gush & Gilbert (1965, 1970) showed that a considerable number of ovulations occur which are not followed by oviposition, particularly when birds are coming into lay. If the ovary exhibits some form of cyclic activity for some time before the onset of laying, the once-weekly sampling may produce patterns which are confounded by the cyclic phenomenon. Although such an explanation may account for the different patterns found, it is unlikely to be the cause of the differences in mean concentration. Some part of these differences may be due to the variation between assays. As is illustrated by the duplicate values obtained for the same plasma sample in two assays (Bird M5), accuracy was reduced with high levels of oestradiol since the extreme portion of the standard curve was employed. Nevertheless, there may be real differences between birds even within the same strain since Heald & Badman (1963) also commented on large between-bird differences in their findings.

The reports of Wood-Gush & Gilbert (1965, 1970) that oviposition is not an accurate indicator of ovulation, particularly with birds coming into lay,
suggests that the time-interval from the peak level of a blood constituent to the time of the first oviposition may be somewhat variable. In further elucidating the process of sexual maturation, it would clearly be preferable to determine metabolic constituents and hormones in the same blood samples.

ACKNOWLEDGMENTS

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REFERENCES


