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Tibial dyschondroplasia, a cartilage abnormality in poultry

B. SAUVEUR, P. MONGIN

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Summary. Tibial dyschondroplasia is characterized by a calcification defect and by the presence of an abnormal mass of unvascularized cartilage in the proximal metaphysis of chick and turkey tibio tarsus. The gross appearance of the lesion shows some similarities with rickets, but the histological figures are quite different, without any modification of the proliferating zone of the epiphyseal plate. The figures of the chemical composition of abnormal cartilage and the blood parameters related to mineral metabolism both remain normal. The abnormal cartilage plug seems to result from the proliferation of prehypertrophic cells in relation to the lack of metaphyseal vascularization.

Tibial dyschondroplasia is under genetic control and can be eliminated after three generations of divergent selection. In the field, its incidence is proportional to the growth rate of the flocks and males are more sensitive than female birds. In unselected birds, tibial dyschondroplasia can be induced by feeding a purified diet containing all known nutrients; corn and soybean meal have preventing properties. The abnormality can not be prevented by any known essential nutrient, such as minerals or vitamins. Metabolic acidosis due to ammonium chloride ingestion or only to an excess of chloride, strongly increases the defect incidence, while growth simultaneously decreases. This adverse effect of chloride can be overcome by the addition of a sufficient level of sodium and potassium into the diet, restoring body weight to a normal value.

Although the tibial dyschondroplasia condition presents a clinical picture quite different from that of rickets, the possible intervention of a vitamin D metabolite is discussed since it is known that metabolic acidosis impairs the renal transformation of 25-(OH)CC into 1,25-(OH)_2CC.

Introduction.

Poultry production was industrialized 40 years ago; since that time, some previously unknown growth disorders have appeared which are mainly ossification defects. The very intensive selection applied to birds (chiefly chicks and turkeys) has increased this problem by enlarging the muscular parts of the body without taking into account skeletal development.

The causes of these bone maturation defects are numerous (Bräunlich, 1972; Sauveur, Roncin and Mongin, 1975). In such simple cases, as for example perosis control by Mn or choline, only one nutritional deficiency is involved. On the other hand, other symptoms do not appear to depend on any dietary deficiency; an example is cartilage abnormalities originally described by Leach and Nesheim (1965a). Their
origin seems to lie in an inherited bone maturation defect aggravated by dietary mineral unbalance of such elements as Na, K and Cl. The purpose of the present paper is to review this type of lesion and to also emphasize the role of acid-base balance in the growth of domesticated birds.

Description of tibial dyschondroplasia.

The cartilage abnormality, tibial dyschondroplasia, is a calcification defect in the proximal metaphysis of the tibio tarsus and began to cause major economic losses in the turkey and broiler industry between 1970 and 1975. This disturbance has been classified as the first non-infectious cause of leg weakness in turkeys and it also represents an important origin of leg problems in chicks and guinea-fowl (in France) as well as in ducks.

The geographical extension of chick dyschondroplasia seems to be unlimited; it was first described at the laboratory level in the United States (Leach and Nesheim, 1965a and b) and then in broiler flocks in Australia and South Africa (Hemsley, 1970), Great Britain (Laursen-Jones, 1970; Siller, 1970; Siller and Duff, 1970), western Canada (Riddel, Howell and Kaye, 1971), United States (Prasad, Hairr and Dallas, 1972), Japan (Itakura, Goto and Fujiwara, 1973) and France (personal observations, 1972, unpublished).

The original name of « cartilage abnormality » given by Leach and Nesheim (1965a) has been progressively replaced by « tibial dyschondroplasia », introduced by Siller (1970). Some other names as « osteochondrodystrophy » (McCapes, 1967) and « cartilage plugs » (Julian, 1973), have been given to cartilage lesions which seemed to be very similar to dyschondroplasia.

Estimates of its frequency in broiler flocks lie between 1 and 5 p. 100; nevertheless, Steinke (1971) had reported an incidence of 33 to 48 p. 100. After examination of 40 broiler-growing farms with a total capacity of 2 million broilers, Prasad, Hairr and Dallas (1972) estimated that the morbidity in affected flocks was 30 to 40 p. 100. It is generally recognized that the incidence is directly related to the growth rate of the flocks.

Tibial dyschondroplasia is not always externally evident. When a flock is severely affected, varied degrees of lameness appear and increase up to slaughtering. Affected birds are reluctant to walk and sit on their hocks; they rise with difficulty when prodded, walk a few steps and immediately resume a sitting position. Such a lameness can be seen after 4 weeks of age in the chicken and after 10 weeks in the turkey. In some cases, the leg bones (mainly the tibio tarsus) show a small curvature but it is not always externally observed. The stifle and hock joints do not swell and the footpads remain normal. The bone strength of affected birds does not generally appear to be less than that of unaffected animals of the same flock.

Bone examination.

Radiographs of the limbs show bilateral symmetric lesions appearing as defined radiolucent areas within the proximal metaphysis of the tibiotarsus and sometimes, of the tarsometatarsus. On longitudinal sections of the proximal end of the bone,
the defect is grossly visible as a white opaque mass (fig. 1), originating under the epiphysis and spreading more or less deeply down through the metaphysis (between 2 and 10 mm at 4 weeks). This abnormal cartilage formation occupies all the bone width in the most severely affected birds, but it is often limited to a narrow lateral band. In older birds (Riddell, Howell and Kaye, 1971), the radiotranslucent area is replaced by a zone of sclerosis. All the authors have noted that there is very little blood vessel tunneling in the abnormal area.

FIG. 1. — Longitudinal sections of proximal ends of tibiotarsus from 3-week old chicks showing dyschondroplasia (Sauveur, 1974, unpublished results).

**Histological examination.**

The gross appearance of the cartilage is similar to that observed in rickets; however, histological examination indicates wide differences. According to the descriptions of Leach and Nesheim (1965a), Siller (1970), Itakura, Goto and Fujiwara (1973) and Riddell (1975), the dyschondroplasic cartilage can be divided into 3 parts, starting from the epiphyseal end:

- a layer of proliferating cartilage of normal constitution which may be slightly thickened or not;
- a layer of pre-hypertrophic or hypertrophic cartilage lacking blood vessels and having a cellular structure similar to that of the normal tissue;
- a mass of abnormal cartilage without vascular invasion in which the morphology and staining properties of the hypertrophic cells are modified; it is often difficult to differentiate the nucleus from the cytoplasm, and the cells do not fill the cavities.

For comparison, it is recalled that rickets, due to vitamin D deficiency in chicks leads to an increase in the width of the proliferating zone (Belanger and Migi-
cowski, 1958) and that EHDP (1) administration results in a proliferation of normal hypertrophic cells (Bisaz et al., 1975). According to Leach and Nesheim (1965a) the lesions of copper deficiency described by Carlton and Henderson (1964) would be quite similar to those in dyschondroplasia.

Chemical examination of abnormal cartilage.

Leach and Nesheim (1965b) noted a small increase of the collagen content of abnormal cartilage. More recently, Lowther et al. (1974) compared the composition of dyschondroplastic cartilage with that of normal epiphyseal growth plate cartilage and of normal articular hyaline cartilage. They concluded that proteoglycan and collagen contents of abnormal cartilage differed considerably from those of articular cartilage but were similar to those of epiphyseal growth plate cartilage of normal and dyschondroplastic birds. Moreover, the proteoglycan subunits isolated from epiphyseal growth plate cartilage of normal birds and from dyschondroplastic cartilage of afflicted animals had similar carbohydrate and protein composition. Lowther et al. (1974) concluded that the abnormal cartilage plug arises from abnormal proliferation of the growth plate cartilage during early development. However, during in vitro incubation, abnormal cartilage slices exhibited a lower rate of proteoglycan synthesis than that recorded with epiphyseal plate cartilage; this might be a factor in the prevention of normal osteogenesis.

TABLE 1

Comparative classification of 4 types of bone abnormalities according to the associated chemical changes
(Leach and Nesheim, 1965b)

<table>
<thead>
<tr>
<th>Type of bone abnormality</th>
<th>Rickets (Vit. D₃ or Ca deficiency)</th>
<th>Rickets (P deficiency)</th>
<th>Perosis (Mn deficiency)</th>
<th>Tibial dyschondroplasia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ca.</td>
<td>——</td>
<td>++</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>P</td>
<td>+</td>
<td>——</td>
<td>——</td>
<td>0</td>
</tr>
<tr>
<td>Alkaline phosphatase</td>
<td>++</td>
<td>0</td>
<td>——</td>
<td>0</td>
</tr>
<tr>
<td>Acid phosphatase</td>
<td>+</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Bone ash</td>
<td>——</td>
<td>——</td>
<td>——</td>
<td>——</td>
</tr>
<tr>
<td>Cartilage</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alkaline phosphatase</td>
<td>0</td>
<td>++</td>
<td>——</td>
<td>0</td>
</tr>
<tr>
<td>Mucopolysaccharide</td>
<td>0</td>
<td>0</td>
<td>——</td>
<td>0</td>
</tr>
<tr>
<td>Collagen</td>
<td>++</td>
<td>0</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>Lyosomal enzymes</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Width of growth plate</td>
<td>++</td>
<td>0</td>
<td>——</td>
<td>0</td>
</tr>
</tbody>
</table>

(1) EHDP = Ethane-1-hydroxy-1, 1-diphosphonate.
Associated symptoms.

Tibial dyschondroplasia is generally characterized by the absence of other pathological signs. Soft tissues surrounding the joints are always normal; bone ash values are similar in afflicted and non-afflicted birds (Leach and Nesheim, 1965a; Riddell, 1975). According to the same authors, calcium, phosphorus, magnesium and phosphatase levels in the blood remain unchanged. Prasad, Hairr and Dallas (1972) also found normal values for blood hematocrit and hemoglobin content. A distinction may thus be easily made between the dyschondroplasic state and some other bone abnormalities such as rickets or perosis (table 1 from Leach and Nesheim, 1965b).

Causes of tibial dyschondroplasia

According to Leach and Nesheim (1965a), it is rather unlikely that abnormal cartilage is the remains of the embryonic type of cartilage cone which can be seen in the tibial diaphysis of the one-day old chick. These authors found that the abnormal cells were morphologically different from those of the diaphyseal cartilage cone, and that fewer lesions were found at 2 weeks than at 4 weeks of age.

Siller (1970) did not agree with these hypotheses and believed that the abnormal cartilage tissue was of embryonic origin, the diaphyseal cartilage cone, which normally disappears within 2 weeks after hatching, remaining intact after this period and proliferating both longitudinally and laterally perhaps because of the defective blood supply. According to Riddell (1975), the failure of vascular invasion from the diaphysis would entail a persistence of prehypertrophic cartilage.

Whatever the exact type of maintained cartilage, Wise and Jennings (1972) also considered that the partial failure of metaphyseal blood supply, a function of which is to calcify and destroy the hypertrophic cartilage, was directly responsible for the accumulation of an uncalcified plug in the metaphysis. According to these authors, this lack of metaphyseal vascularization itself would result from abnormal pressures applied to bone epiphysis due to excessive growth rate. These authors furthermore reported that lesions of the lower ulnar were also known in dogs and pigs and that these calcification defects were thought to be caused by abnormal compression and interruption of blood supply (Riser and Shirer, 1965; Vaughan, 1971). Such a structural explanation of tibial dyschondroplasia is in agreement with the observed interrelationships between dyschondroplasia incidence and the growth rate of broiler flocks, and with the predominance of the lesion in male as compared to female birds. However, this hypothesis does not explain several genetic and nutritional observations discussed below.

Genetic investigations.

The original work of Leach and Nesheim (1965a) has clearly demonstrated that the susceptibility of chickens to tibial dyschondroplasia is under genetic control. The authors have developed two divergent strains in which the incidences were 40 to 50 and 15 p. 100, respectively, after only one generation; 6 years later Leach (1971) announced values of 80 to 90 vs 0 p. 100. It must also be noted that, contrary to what
is seen in field observation, the low-incidence strain was heavier than the high-incidence strain. By divergent selection for three generations, Riddell (1975, 1976) also recently obtained two populations of broilers with dyschondroplasia incidences equal to 51 and 0 per 100, respectively. This author noted that other skeletal defects as spondylolisthesis and twisted legs were unrelated to tibial dyschondroplasia, an observation already made by Mongin and Sauveur (1973). Steinke (1971), comparing two strains of turkeys, reported that the heavier strain had a higher incidence of tibial dyschondroplasia than the smaller strain; the cross of the two strains was intermediate in weight and in amount of abnormal cartilage. The same observation has been made in chickens.

All the studies showed greater susceptibility of male chicks as compared to females. However, at sufficient levels to alter secondary sex characteristics, neither testosterone propionate nor diethylstilbestrol had any significant effect upon the occurrence of cartilage abnormality in male or female chicks (Leach and Nesheim, 1965a). It is thus more probable that the sex effect was due to the different growth rate.

**Nutritional investigations.**

In the first study of Leach and Nesheim (1965a), a 31 per 100 incidence of tibial dyschondroplasia was found in chicks fed a *purified control diet* containing glucose (58.5 per 100), isolated soybean protein (27.0 per 100), vitamin, mineral and amino acid mixtures, and believed to be adequate in all known nutrients; the defect was absent in chicks fed a commercial starter diet. Mixtures of chick starter with basal diet indicated that 50 to 100 per 100 chick starter was required for complete prevention of the abnormality (table 2). During a study in which the ingredients of usual commercial chick rations were added to the purified diet, only corn and soybean meal had some dyschondroplasia-preventing properties. The same study (Leach and Nesheim 1965a) showed that the abnormality was not due to a deficiency or an excess of the following nutrients in the purified diet:

- minerals: Ca, P, Mg, Fe, Cu, Zn and Se;

<table>
<thead>
<tr>
<th>Purified basal diet (p. 100)</th>
<th>Chick starter (commercial diet) (p. 100)</th>
<th>Incidence of tibial dyschondroplasia (p. 100)</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>0</td>
<td>60 (20) *</td>
</tr>
<tr>
<td>75</td>
<td>25</td>
<td>33 (21)</td>
</tr>
<tr>
<td>50</td>
<td>50</td>
<td>14 (21)</td>
</tr>
<tr>
<td>0</td>
<td>100</td>
<td>0 (20)</td>
</tr>
</tbody>
</table>

* Number of male chicks.
— vitamins: D₃, D₂, niacin, pyridoxine and inositol. Thus, although tibial dyschondroplasia has some common visible traits with rickets or copper deficiency, its causes appear to be quite different.

Similar observations were made by Steinke (1971) in turkeys, and the following treatments were without any preventing effect:
- enhanced levels of Ca, P, Zn, Cu, Mo; addition of EDTA;
- normal levels multiplied 2 to 10 times of vitamins A, D₃, E, K, C, B₆, B₁₂, pantothenic acid, niacin, choline, biotin, folacin, thiamin;
- utilization of 4 different commercial sources of vitamin D₃, dihydrotachysterol and lumisterol;
- some treatments such as restricted feeding, aureomycin supplementation.

These results demonstrated clearly that tibial dyschondroplasia cannot be prevented by any known nutrient. Two hypotheses were thus proposed: either an unidentified growth factor was involved, or the nutrients needed to be balanced. The second proposition appeared to be the most interesting.

Between 1970 and 1972, Mongin, Leach and Nesheim together conducted a series of experiments showing that blood acid-base balance could play a role in dyschondroplasia; metabolic acidosis induced by feeding ammonium chloride increased dyschondroplasia incidence, even in birds fed a commercial corn-soybean diet. This fact is illustrated in table 3 (Leach and Nesheim, 1972). The incidence of dietary dyschondroplasia could be reduced when H⁺ ion was accompanied by anions such as sulfate or acetate, as shown in table 4 (Mongin, Leach and Nesheim, 1970, unpublished results). These anions reduced acidosis severity in two different ways: the sulfate ion, poorly reabsorbable by the renal tubule, increases urinary H⁺ excretion as demonstrated in laying hens by Sauveur (1969), and the metabolisable

\[
\begin{array}{|c|c|c|c|c|}
\hline
\text{Diet} & \text{Added NH₄Cl (p. 100)} & \text{4 week weight (g)} & \text{Incidence of dyschondroplasia (p. 100)} & \text{Blood composition} \\
& \text{p. 100} & \text{pH} & \text{HCO₃⁻ (mEq. L⁻¹)} \\
\hline
\text{Purified} & 0.00 & 657 & 20 & 7.18 & 18.8 \\
& 0.73 & 575 & 71 & 7.11 & 14.9 \\
& 1.46 & 508 & 76 & 7.08 & 12.2 \\
\text{Corn-Soybean} & 0.00 & 671 & 0 & 7.23 & 21.9 \\
& 0.73 & 667 & 9 & 7.23 & 21.2 \\
& 1.46 & 614 & 33 & 7.21 & 18.6 \\
\hline
\end{array}
\]

\[\text{H⁺-induced dyschondroplasia could be reduced when H⁺ ion was accompanied by anions such as sulfate or acetate, as shown in table 4 (Mongin, Leach and Nesheim, 1970, unpublished results). These anions reduced acidosis severity in two different ways: the sulfate ion, poorly reabsorbable by the renal tubule, increases urinary H⁺ excretion as demonstrated in laying hens by Sauveur (1969), and the metabolisable}\]
acetate anion consumes the $H^+$ ion during its catabolism and annuls $NH_4^+$ loading (Richet, Ardaillou and Amiel, 1966).

The results of Leach and Nesheim (1972) also showed that, without any $H^+$ addition to the diet, distribution of a high chloride mineral mixture ($Cl = 0.84 \text{ p. 100}$ of the diet) was sufficient to induce a very high incidence of dyschondroplasia (43 p. 100 in unselected birds and 91 p. 100 in a selected strain fed a purified diet). Similar results have been reported recently by Riddell (1975, 1976) with diets containing 0.75 p. 100 chloride. The problem of chloride effect upon cartilage abnormality has been studied in more detail in our laboratory. As indicated in table 5 (Sauveur and Mongin, 1974), the plasma bicarbonate decreased and dyschondroplasia incidence increased when an excess of chloride was not balanced by cation loading (Na and K). On the contrary, the effect of high chloride level could be strongly reduced when cations were added into the diet at a sufficient level. It also appeared that an excess of sodium relative to potassium raised dyschondroplasia incidence; this fact was particularly obvious with the highest addition of chloride. In every case, there was good parallelism between abnormality incidence and the metabolic acidosis severity recorded.

**TABLE 4**

*Effect of anions accompanying dietary $H^+$ on tibial dyschondroplasia incidence*  
(Mongin, Leach and Nesheim, 1970, unpublished results)

<table>
<thead>
<tr>
<th></th>
<th>4 week weight (g)</th>
<th>Incidence of dyschondroplasia (p. 100)</th>
<th>Blood composition</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>$pH$</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Purified</td>
<td>661</td>
<td>14.7</td>
<td>7.25</td>
</tr>
<tr>
<td>$+ \text{CaCl}_2 \cdot 2H_2O$ (1 p. 100)</td>
<td>589</td>
<td>28.1</td>
<td>7.16</td>
</tr>
<tr>
<td>$+ \text{NH}_4\text{Cl}$ (0.73 p. 100)</td>
<td>581</td>
<td>25.6</td>
<td>7.17</td>
</tr>
<tr>
<td>$+ (\text{NH}_4)_2\text{SO}_4$ (0.9 p. 100)</td>
<td>578</td>
<td>18.8</td>
<td>7.21</td>
</tr>
<tr>
<td>$+ \text{NH}_4$ Acetate (1.05 p. 100)</td>
<td>658</td>
<td>6.1</td>
<td>7.27</td>
</tr>
</tbody>
</table>

The results of Leach and Nesheim (1972) also showed that, without any $H^+$ addition to the diet, distribution of a high chloride mineral mixture ($Cl = 0.84 \text{ p. 100}$ of the diet) was sufficient to induce a very high incidence of dyschondroplasia (43 p. 100 in unselected birds and 91 p. 100 in a selected strain fed a purified diet). Similar results have been reported recently by Riddell (1975, 1976) with diets containing 0.75 p. 100 chloride. The problem of chloride effect upon cartilage abnormality has been studied in more detail in our laboratory. As indicated in table 5 (Sauveur and Mongin, 1974), the plasma bicarbonate decreased and dyschondroplasia incidence increased when an excess of chloride was not balanced by cation loading (Na and K). On the contrary, the effect of high chloride level could be strongly reduced when cations were added into the diet at a sufficient level. It also appeared that an excess of sodium relative to potassium raised dyschondroplasia incidence; this fact was particularly obvious with the highest addition of chloride. In every case, there was good parallelism between abnormality incidence and the metabolic acidosis severity recorded.

**TABLE 5**

*Effect of (Na $\div$ K) and Cl levels (mEq/100 g) and Na/K ratios in the diet on tibial dyschondroplasia incidence (p. 100)*  
(Sauveur and Mongin, 1974)

<table>
<thead>
<tr>
<th>Na/K</th>
<th>Cl level (Na $\div$ K) level</th>
<th>10</th>
<th>25</th>
<th>40</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>20</td>
<td>35</td>
<td>50</td>
<td>20</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>16.6</td>
</tr>
<tr>
<td>2.5</td>
<td>10</td>
<td>7.6</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>0.4</td>
<td>7.1</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>
It may be asked why metabolic acidosis impairs the normal cartilage maturation. It is well known that vitamin D₃ has to be transformed into 1,25 dihydroxycholecalciferol in the kidney before it becomes metabolically active (De Luca and Schnoes, 1976). Sauveur et al. (1977) have shown both in vitro and in vivo that metabolic acidosis reduced the renal conversion of 25-(OH)₃CC into 1,25-(OH)₂CC by about 40 p. 100 in rachitic chicks. The same result has been reported in rats (Lee, Russell and Avioli, 1977). It could then be supposed that vitamin D₃ metabolism was involved in dyschondroplasia, but it has been said previously that the cartilage looks different in rickets and dyschondroplasia, and it would be surprising that a defect of cholecalciferol metabolism acted only on the tibial metaphysis. Several laboratories are now investigating the direct effect of 1,25-(OH)₂CC or 1α-(OH)₂CC on tibial dyschondroplasia.

The only conclusion which can be drawn today is therefore that of Leach and Nesheim (1972) : "Tibial dyschondroplasia is the result of an inherited physiological defect, the expression of which is under dietary control". We have some data on this dietary control (role of chloride, anion/cation ratio), but no hypothesis which would yet explain all the recorded observations.

**Relationships between mineral nutrition, cartilage abnormalities and body growth**

The studies reported here are interesting, not only in relation to cartilage abnormality, but also because mineral and acid-base balance play a misappreciated role in the weight gain of domestic birds. It can be seen in table 3 that the simple utilization of a purified diet was sufficient to induce metabolic acidosis and a slight reduction of growth (Leach and Nesheim, 1972).

The equilibrium between chloride and sodium plus potassium levels in the diet is also important. Nesheim et al. (1964) have demonstrated that an excess of chloride depressed growth response of chicks unless this anion was balanced with equimolar levels of Na or K supplied by a metabolisable anion. This fact was confirmed by Mongin and Sauveur (1973) using carbonate and chloride mixtures of Na and K with a constant Na/K ratio equal to the unity. When Na and K are not used at the same level (expressed in milliequivalents), chicks better tolerate an excess of potassium than an excess of sodium (Nesheim et al., 1974 ; Sauveur and Mongin, 1974).

Several authors have tried to determine the exact anion/cation ratio compatible with an optimal growth ; according to Melliere and Forbes (1966), it would be in the range of 1.2 to 1.8 (on a mEq basis). More recently, Hurwitz et al. (1973, 1974) found that the optimal Na/Cl ratio (w/w) was 1, i.e. 1.5 on a mEq basis. An unpublished work of Mongin and Sauveur (1973) including 24 different semisynthetic diets (3 chloride, 2 sodium and 4 potassium levels) showed (fig. 2) that the best way to get a simple relationship was to use the amount (Na + K — Cl), expressed in mEq/100 g of diet, as a variable. Under such conditions, all the experimental values of weight gain were distributed around a parabolic curve, the maximum of which was obtained for a value of (Na + K — Cl) close to 25 mEq/100 g of diet. The same value was found again under practical conditions with 1 200 8-week old broilers fed a classical
cornsoybean diet complemented with 12 different mixtures of Na, K and Cl (Sauveur and Mongin, 1974, unpublished).

In conclusion, there are strict interrelationships between Na, K and Cl levels in the diet of broiler chicks, and it would be incorrect to fix the requirement for one of these minerals without knowing the levels of the other two. These observations can be particularly important when birds are fed amino acid hydrochloride mixtures.

Résumé. La dyschondroplasie tibiale est un défaut d’ossification des têtes tibiales se traduisant par la persistance d’une masse cartilagineuse non vascularisée dans la métaphyse proximale du tibiotarse des poulets et des dindons. L’apparence globale de la lésion pourrait faire penser à du rachitisme ; cependant ni l’étude histologique ni la symptomatologie générale ne confirme cette possibilité : au niveau histologique, la zone de prolifération de la plaque épiphysaire reste normale alors qu’elle est élargie dans le rachitisme ; par ailleurs ni la composition chimique du cartilage (glycoprotéines et collagène), ni les paramètres sanguins liés au métabolisme phospho-calcique (Ca, P, Mg, phosphatase) ne...
sont significativement modifiés. L'accumulation anormale de cartilage paraît résulter d'une prolifération de cellules à l'état pré-hypertrophié en relation avec l'absence de vascularisation métaphysaire.

La sensibilité à la dyschondroplasie tibiale paraît hautement héritable et il est possible d'éliminer totalement le trouble d'un troupeau après trois générations de sélection divergente. Sur le terrain, l'incidence est directement proportionnelle à la vitesse de croissance des troupeaux et les mâles sont toujours plus sensibles que les femelles. On peut, chez des oiseaux non sélectionnés, provoquer l'apparition de la maladie en utilisant des régimes purifiés contenant cependant tous les nutriments connus. Aucun élément minéral (macro- ou oligo-constituant) ni aucune vitamine n'exerce d'effet curatif contrairement au maïs et au tourteau de soja ! Une acidose métabolique due à l'ingestion de chlorure d'ammonium ou à un simple excès de chlore alimentaire, augmente fortement l'incidence de la maladie (alors que le poids vif de l'animal diminue simultanément). Cet effet nocif du chlore est supprimé par un apport simultané de sodium et de potassium accompagnés d'un anion métabolisable.

Bien que la dyschondroplasie du poulet paraisse donc relever de mécanismes différents de ceux de rachitisme, l'intervention éventuelle d'un trouble du métabolisme de la vitamine D est évoquée puisque l'on sait que l'acidose métabolique, en tant que telle, altère la synthèse rénale de $1,25-(OH)_2$CC à partir du $25-(OH)_2$CC.

References


