Perspectives in mineral nutrition

By T. G. TAYLOR, Department of Physiology & Biochemistry, University of Southampton, Southampton SO9 3TU

The last symposium on poultry nutrition organized by the Nutrition Society was held in Edinburgh 13 years ago and in this paper some of the more important advances that have been made in the mineral nutrition of poultry during this period will be reviewed. This period coincides roughly with the period between the establishment by the Agricultural Research Council of the first Working Party on Nutrient Requirements of Poultry (whose report was published in 1963) and the completion of the work of its successor (Agricultural Research Council, 1975).

Trace elements

One of the most exciting developments in mineral nutrition during the past 13 years has been the extension of the list of elements known, with certainty, to be essential for poultry to include nickel, tin, vanadium, silicon and fluorine (see Schwarz (1974) for review and references).

Requirements for these newer trace elements are incredibly minute and a deficiency of most of them can only be demonstrated in animals maintained in isolators and given highly purified diets and it is not likely, therefore, that deficiencies will arise under practical conditions. It would be unwise, however, to dismiss this possibility completely. A deficiency of chromium, on the other hand, can certainly be demonstrated without the use of isolators and it may therefore turn out to be of practical significance in poultry nutrition as it is thought to be in human nutrition (Schroeder, 1968).

The status of selenium has changed somewhat since 1961; at that time a requirement for Se that could not be provided for by vitamin E had not been demonstrated. The identification of a Se-deficiency condition in Japanese quail and in chicks (Thompson & Scott, 1967) in the presence of an abundance of vitamin E established that Se was an essential trace element in its own right, and it may even prove to be the senior member of the Se-vitamin E partnership. Diplock & Lucy (1973) have suggested that a major role of vitamin E may be to protect the iron-selenide active centres of certain membrane-bound proteins that play a key role in electron transport in mitochondria and possibly in the endoplasmic reticulum.

Recommended requirements for a number of the ‘older’ trace elements have increased in recent years. The Agricultural Research Council (1963) recommended requirement for zinc for young chicks was 15 mg/kg diet, but it is now recognized
that the true requirement is much higher than this in practical diets, but that under most husbandry conditions substantial amounts of Zn are derived from the environment, particularly from drinking troughs. High concentrations of calcium and phytate reduce the availability of dietary Zn (O'Dell, Yohe & Savage, 1964) but there are other factors, still unknown, that influence the availability of Zn (Turk & Lease, 1962). The true requirement for chicks appears to be at least 40 mg/kg diet (Ziegler, Leach, Norris & Scott, 1961) and for turkey poults 64 mg/kg (Sullivan, 1961) when soya-bean protein is used as a protein source.

The requirements of chicks for Fe appear now to be greater than the 40 mg/kg diet suggested by Hill & Matrone (1961) and Davis, Norris & Kratzer (1968) conclude that 75–80 mg/kg are required. As with many other nutrients, observed requirements vary somewhat according to the criteria used: with Fe, growth, haemoglobin concentration and packed cell volume are the normal criteria employed.

**Calcium, phosphorus and vitamin D Metabolic studies**

Advances in knowledge concerning the metabolism of Ca have been truly spectacular since 1961, beginning with the discovery of calcitonin (CT) (Copp, Davidson & Cheney, 1961). Subsequent advances centred on vitamin D: on its mode of action and metabolism and on the relationships between parathyroid hormone (PTH), CT and the vitamin; much of this fundamental work was carried out in the chick.

The first indication of the mechanism of action of vitamin D in increasing Ca absorption from the gut came with the discovery that Actinomycin D, an inhibitor of protein synthesis, prevents the increase in Ca absorption in rachitic animals treated with vitamin D (Norman, 1965; Zull, Czarnowska-Misztal & De Luca, 1965). The following year Wasserman & Taylor (1966) demonstrated the presence of a vitamin D-dependent Ca-binding protein (CaBP) in the duodenum of normal and vitamin D-treated chicks and a similar protein was subsequently identified in kidney (Taylor & Wasserman, 1967) and in the shell gland (Corradino, Wasserman, Pubols & Chang, 1968).

During the next few years a number of biologically active metabolites of cholecalciferol were identified and we now know that the normal metabolism of cholecalciferol involves, first, hydroxylation by the liver to give 25-hydroxycholecalciferol (25-HCC) (Blunt, De Luca & Schnoes, 1968), the main form in which vitamin D is transported in the blood. Subsequently this compound is hydroxylated by the kidney to give 1,25-dihydroxycholecalciferol (1,25-DHCC) (Lawson, Fraser, Kodicek, Morris & Williams, 1971). This compound is the most potent derivative of vitamin D known and it is now considered to be the physiologically active form of the vitamin, acting in the control of protein synthesis in target tissues at the transcription stage in a manner similar to the action of steroid hormones.

The subcellular localization of CaBP in the intestinal mucosa is still not known for certain, nor is the mechanism by which it promotes Ca transport understood. We
do know, however, that in most physiological situations in the chick and in the rat there is a very close relationship between the concentration of CaBP in intestinal tissue and the rate of Ca absorption both in vivo and in vitro. Thus, for example, CaBP increases in response to low-Ca diets and decreases when high-Ca diets are given (Taylor & Wasserman, 1969). It is now clear that 1,25-DHCC is the major factor controlling Ca absorption by the gut of chicks (by controlling CaBP synthesis) and the amount of this active metabolite in the circulation is controlled in vitamin D-replete animals by changes in the activity of the 1-hydroxylase in the kidney. In normocalcaemic and hypercalcaemic states the main metabolite of 25-HCC in the kidney is 24,25-DHCC, while in hypocalcaemic conditions 1,25-DHCC predominates (Holick, Schnoes, De Luca, Gray, Boyle & Suda, 1972). In the rat there would also appear to be a relationship between plasma inorganic P concentration and the metabolism of 25-HCC by the kidney: above a concentration of 2.6 mmol P/l, 24,25-DHCC is the major product, while at concentrations below this 1,25-DHCC is largely formed (Tanaka & De Luca, 1973).

There is still some uncertainty concerning the relative importance of PTH, Ca and inorganic phosphate in controlling the activity of the kidney 1-hydroxylase, but the results of in vivo experiments in the chick strongly suggest that PTH is primarily responsible for this control; a high concentration of the hormone is associated with high enzyme activity (Fraser & Kodicek, 1973). Rasmussen, Wong, Bikle & Goodman (1972) showed that PTH and cyclic AMP stimulated the conversion of 25-HCC to 1,25-DHCC by isolated chick kidney tubules and that CT inhibited the conversion.

Practical considerations

(a) Chicks and poults. It has been recognized for many years that minimum requirements for Ca and P vary inversely according to the amount of vitamin D in the diet. Thus, for example, Waldroup, Stearns, Ammerman & Harms (1965) observed that when chicks were fed on a diet containing 10 g Ca/kg, requirements for cholecalciferol for maximum growth and bone mineralization were no more than 5 μg/kg, but that when the Ca concentration was reduced to 5 g/kg diet, requirements for cholecalciferol increased to 40 μg/kg. At this dietary concentration of cholecalciferol, growth was as good as at the higher Ca concentration but bone-ash percentage was greatly reduced and even at 400 μg/kg bone mineralization was poor. It is now possible to offer an explanation for these observations in the light of recent developments in vitamin D metabolism. When chicks are given a low-Ca diet the plasma Ca tends to fall, PTH secretion is stimulated and synthesis of 1,25-DHCC is increased, stimulating CaBP synthesis in the intestinal mucosa. Provided the dietary Ca is only slightly reduced and the concentration of vitamin D in the diet is adequate, the increase in the CaBP in the intestines will stimulate Ca absorption sufficiently to allow normal growth and calcification of the skeleton. With diets seriously deficient in Ca this element becomes limiting for bone calcification and it may even become limiting for growth, in spite of an abundant supply of vitamin D.

Any increase in the rate of synthesis of 1,25-DHCC (which has a very short half-life
in the plasma) increases the dietary requirement for vitamin D by increasing the utilization of 25-HCC. Vitamin D is required for growth in addition to its requirement for Ca absorption and for bone metabolism (striated muscle is one of the target tissues for 1,25-DHCC). If, therefore, supplies of vitamin D are not increased when the dietary Ca is lowered, growth will be reduced.

Turkey poults are particularly sensitive to deficiencies, relative or absolute, of Ca and P in the diet. Neagle, Blaylock & Goihl (1968) observed a mean live weight of 554 g at 4 weeks of age in poults given diets containing 8 g Ca and 6 g P/kg and varying amounts of vitamin D, all considered to be adequate for normal growth. When the dietary Ca was increased to 16 g/kg the mean weight was 484 g for the same concentration of P and the same amounts of vitamin D. Excess P depressed growth in a similar manner when diets marginally deficient in Ca (6 g/kg) were given, unless the concentration of dietary vitamin D was increased.

In the past we have sought to explain the inter-relationship between dietary concentrations of Ca, P and vitamin D largely in terms of the availability of the minerals and the formation in the gut of insoluble complexes, especially phytates (Taylor, 1965), but it is clear that these inter-relationships must now be re-evaluated experimentally in the light of the newer knowledge of vitamin D metabolism. The turkey poult would seem to be an ideal subject for this re-evaluation because of its sensitivity to changes in dietary Ca and P and because of its high requirement for vitamin D. Is growth depression on diets with high or low Ca:P ratios associated with reduced 1,25-DHCC synthesis caused in the former instance by a depression in PTH secretion or an increased CT secretion and in the latter instance by an elevation in the plasma inorganic P concentration?

With the scarcity and high cost of P supplements it is particularly important that the fullest use should be made of the P in cereals and protein concentrates and that the margin of safety allowed in dietary formulations should be minimal. This emphasizes the need for the requirement for P for each class of stock to be known as accurately as possible. Maximum availability of phytate P is achieved when dietary Ca concentrations are low, so it is important that requirements for Ca should not be exceeded if the aim is to utilize dietary P with the maximum efficiency. The Agricultural Research Council (1963) estimated requirement of Ca for turkey poults is 15 g/kg, but more recent reports (Nelson, Jensen & McGinnis, 1963; Sullivan & Kingan, 1963; Neagle et al. 1968) suggest that 9 g/kg is probably adequate.

(b) Laying hens. Results of metabolic and nutritional studies with chicks and poults are consistent with the idea that the 1,25-DHCC-dependent CaBP plays a key role in controlling Ca absorption by the intestines, but the situation in the laying bird is by no means so clear-cut. The two major phenomena that require to be explained are as follows. (1) Ca absorption increases greatly during the 10–14 d period prior to the laying of the first egg under the combined influence of oestrogen and androgen (Common, Rutledge & Hale, 1948). (Oestrogen and androgen alone have only the smallest effect on Ca absorption.) (2) Ca absorption is about twice as great on egg-forming as on non-egg-forming days (Taylor & Kirkley, 1967).

No information has been published on the CaBP in the intestines during the
immediate pre-laying period during which Ca absorption is substantially increased, but Bar & Hurwitz (1972) observed only a slight increase in duodenal CaBP in pullets killed 'at the start of visible maturation' compared with birds 4 months of age killed 10 d earlier. It is unlikely, however, that birds showing signs of sexual maturation had in fact reached the stage at which increased Ca absorption was occurring. In birds killed after laying their first egg, CaBP activity had increased significantly compared with immature birds and, 58 d later, the activity was more than double that observed at the onset of reproduction. The activity declined from this time but 98 d after the onset of lay it was still higher than at the time of the first oviposition. Towards the end of the first laying year the CaBP activity had fallen to the level observed at the start of lay. Arrest of egg production by dosing birds with the drug Nicarbazin resulted in a reduction of CaBP to non-laying levels after 9 d of treatment, but on withdrawing the drug from the diet CaBP levels were restored within a week of resumption of production.

No increase in CaBP activity was observed in the duodenum of hens given a low-Ca diet for 5 d (Hurwitz & Bar, 1969) but after 16 d on a diet containing 17 g Ca/kg a slightly increased activity was observed, and after 32 d the increase was substantial (Bar & Hurwitz, 1972).

No changes have been reported in CaBP during the egg cycle in birds laying regularly and since it is assumed that the CaBP synthesized in any particular mucosal cell remains for the life span of that cell (approx. 3 d) it is difficult to envisage a role for CaBP in the short-term regulation of Ca absorption that occurs in relation to shell calcification. Nevertheless, differences in the activity of 25-HCC 1-hydroxylase in the kidney of Japanese quail during the egg cycle have been reported by Kenny, Lamb, David & Losty (1974). Substantially higher activities of this enzyme were observed when shell calcification was in progress than when the shell gland was empty. It seems probable that enhanced synthesis of 1,25-DHCC during shell formation is more likely to be related to the requirements for bone resorption than for CaBP synthesis by the intestines.

The available evidence suggests that there are two distinct mechanisms involved in the intestinal absorption of Ca in the laying hen, one CaBP-dependent and one independent of CaBP. It would appear that CaBP is largely responsible for raising the basal rate of Ca absorption from the non-laying to the laying rate (even on non-egg-forming days Ca absorption is substantially higher than in non-laying pullets or hens). Whether or not the duodenum alone is involved in this action is not known, since detailed studies of CaBP activities in the jejenum and ileum of laying hens do not appear to have been published. Nor is the stimulus for increasing the production of 1,25-DHCC in laying compared with non-laying pullets known, but it may well be PTH. (It is unlikely to be oestrogens or androgens since these hormones are thought to be secreted normally in Nicarbazin-treated birds and, furthermore, Wasserman & Taylor (1968) have demonstrated that CaBP is not induced by diethylstilboestrol.)

The nature of the second mechanism for Ca absorption, independent of CaBP, is quite obscure, but one question that immediately arises is: 'Is it also dependent on
vitamin D?' Hens suffering from a chronic deficiency of vitamin D can certainly absorb Ca: although they lay few eggs, these have normal shells, and the deficient birds often possess abnormally large amounts of medullary bone (Taylor, 1970).

In spite of the rapid advances that have been made on the basic mechanisms controlling Ca absorption during the last few years, there is clearly a great deal still to be learned, particularly in the laying bird. The most pressing question that needs to be answered concerns the mechanism of action of CaBP. We also need to know what relationship, if any, exists between intestinal CaBP activity, shell thickness and stage of production. Which is the more important for shell thickness, CaBP activity or the 'second mechanism' for Ca transport? Do both mechanisms decline during the latter stages of the laying year?

Let us hope that these questions and the ones noted earlier in this paper will be answered by the time the Society next arranges a symposium on poultry nutrition in Edinburgh.

REFERENCES


