Various Levels of Calcium and Phosphorus Diets in Response to 1, 25-Dihydroxycholecalciferol in Laying Hens

M. Shivazad¹*, A. B. Carlos², H. M. Edwards, JR². and M. Zaghari¹

ABSTRACT

The effect of 1,25-dihydroxycholecalciferol [1,25-(OH)₂ D₃ ] supplementation on laying hen diets was evaluated using diets high and low in Ca (3.0% and 1.8%) and high and low in P (0.52% and 0.33%). Since two levels of 1,25-(OH)₂ D₃ supplementation (0 and 5 µg/kg diet) were applied, the experiment was a 2 X 2 X 2 factorial arrangement in a completely randomized design using nine individually housed hens per each 8 treatments (72 in total). The hens were fed the diets for 28 days and records were kept on body weight, egg production, feed intake, egg weight, and egg specific gravity. At the end of the experiment, the hens were bled for plasma Ca and P determination and after being euthanased the left tibia removed for bone ash measurement. The corn-soybean meal based diet contained 0.1% Cr₂O₃ as an indicator for determining Ca, P and phytate phosphorus retention at 14 and 28 days. 1,25-Dihydroxycholecalciferol supplementation had no effect on hen weight, egg production or feed intake. However, large increases in egg specific gravity were obtained when 1,25-(OH)₂ D₃ was fed to hens receiving the low Ca diet. All of the treatments had significant effects on bone ash. The greatest effect of 1,25-(OH)₂ D₃ on bone ash was obtained in those hens fed the high Ca and P diet that was then supplemented with 1,25-(OH)₂ D₃ (49.3% vs 53.9% bone ash). At day 14, the high Ca diets decreased phytate P retention while at day 28 the high P diets decreased phytate P retention.

Keywords: Calcium, 1, 25-dihydroxycholecalciferol, Phytate phosphorus, Phosphorus.

INTRODUCTION

When laying hens were fed a corn-soybean meal based diet, they utilized only 8% of the phytate P present in that diet as measured by phytate disappearance (Nelson, 1976). This finding is in agreement with the poor utilization of phytate P by laying hens reported by Gillis et al. (1953). Sebastian et al. (1998) listed the following as factors affecting phytate phosphorus utilization: dietary calcium and phosphorus concentration, dietary vitamin D₃ concentration, age of bird, phytase activity of dietary ingredients, fiber and genotype. Qian et al (1997) determined that calcium retention ranged between 42% and 67%, depending on the calcium: total phosphorus ratio, the addition of 66 or 666 µg kg⁻¹ diet vitamin D₃, and the addition of 0-900 phytase units kg⁻¹ diet. No critical studies have been conducted on the effect of dietary Ca and P levels on the utilization of phytate P by laying hens. Recently it was found that the addition of the vitamin D₃ derivative 1,25-dihydroxycholecalciferol (1, 25-(OH)₂ D₃ ) to a corn-soybean meal diet for broilers stimulated phytate P utilization (Edward, 1993).

The experiment reported in this paper was conducted to determine whether supplementation of corn-soybean meal laying diets with 5 mg of a 1, 25-(OH)₂ D₃ / kg diet would stimulate natural phytate utilization at adequate and low levels of dietary Ca and P.

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MATERIAL AND METHODS

Seventy-two Single Comb White Leghorn hens were selected from among 108 sixty week-old hens during a 10 day pre-experiment period. The hens were allocated on the basis of their egg production in 24 pens with three individual cages equalizing average egg production within the pens. The corn-soybean meal based diet presented in Table 1 was utilized during a 28 day period. All substitutions were made at the expense of corn. Limestone (39%Ca) was utilized as a source of calcium. Commercial dicalcium phosphate (analyzed 23.5%Ca and 19%P) was used as a P and Ca source. Two levels of Ca (3.0% and 1.8%) with two levels of total P (0.52% and 0.33%) and two levels of 1,25-(OH)2 D3 (0 and 5 µg/kg) were used in a 2 X 2 X 2 factorial arrangement in a completely randomized design. The hens were allowed to consume feed and water ad libitum. Chromic oxide added to the all treatment diets as an indicator. Collection of excreta was carried out twice every two weeks for 48 hours during the 28 days of the experiment. This was accomplished by setting individual trays under each hen. Dried feed and excreta samples were analyzed for Ca, P and phytate phosphorus by modified HPLC method of Bos et al (1991) and the percent retention of Ca, P and phytate phosphorus was calculated. At the end of the experiment, the hens were individually weighed and bled by heart puncture for analyzing plasma Ca and P levels. They were then euthanased by cervical dislocation and their left tibia was removed for bone ash determination based on fat-free by method (Association of Official Agricultural Chemists, 1955).

Egg production was recorded daily and feed consumption was measured at the end of the experiment. Egg specific gravity based on the flooding method and egg weight were determined weekly. From these data, average daily feed consumption per hen and, feed required per unit of egg weight were calculated. All statistical analyses were conducted on pen average. Analyses of data were computed using the SAS analysis of variance procedure (Helwing and Council, 1979). Main effects were separated by Duncan’s multiple range test.

RESULTS

There were no significant treatment effects on hen body weight (BW) or feed intake. These data are not presented. Only the Ca level caused a significant effect on feed/g egg (g/g). The birds receiving the high Ca diet required 2.0 g feed/g egg and those receiving the low Ca diets consumed 2.4 g feed/g egg, these data are not presented. Data on weekly egg production, egg weight and egg specific gravity were analyzed using the week as a continuous variable. The egg production data showed significant effects for both diet and week with no interaction, indicating that the significant effect of the low Ca diet in reducing egg production was just an acceleration of the general decline in egg production by all treatments during the experiment (Table 2). Neither the level of P nor the addition to the diet of the 5 mg/kg of

Table 1. Composition of basal ration

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>%</th>
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<tbody>
<tr>
<td>Ground yellow corn</td>
<td>75.23</td>
</tr>
<tr>
<td>Soybean meal</td>
<td>22.00</td>
</tr>
<tr>
<td>Poultry oil</td>
<td>2.00</td>
</tr>
<tr>
<td>Salt (sodium chloride)</td>
<td>0.30</td>
</tr>
<tr>
<td>Vitamin premix*</td>
<td>0.25</td>
</tr>
<tr>
<td>Trace mineral premix*</td>
<td>0.05</td>
</tr>
<tr>
<td>DL-methionine</td>
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<tr>
<td>Chromic oxide</td>
<td>0.10</td>
</tr>
</tbody>
</table>

* Vitamin premix provides (per kg of diet): vitamin A, 5, 500 IU; vitamin D3, 1,100 ICU; vitamin E, 11 IU; riboflavin, 4.4 mg; calcium pantothenate, 12 mg; nicotinic acid, 44 mg; choline cl, 220 mg; vitamin B12, 9 µg; vitamin B6, 3.0 mg; menadione, 1.1 mg (as menadione sodium bisulfite); folic acid, 3 mg; d-biotin, 3 mg; thiamine, 2.2 mg (as thiamine mononitrate); ethoxyquin, 125 mg.

* Trace mineral premix provides (PPm of diet): MnO2, 111; Zn no, 75; FeSO4.7H2O, 100; CaSO4.14; (IO3)2, 7.5; Ca 7.5; FeCO3, 41.5.
Table 2. Effect of dietary 1,25-dihydroxycholecalciferol (1,25-(OH)₂D₃), Ca and P on egg production, Egg weight and specific gravity, plasma Ca and P, bone ash and phytate P retention by laying hens

<table>
<thead>
<tr>
<th>Ca (%)</th>
<th>Dietary Content (%)</th>
<th>P (µg/kg)</th>
<th>1,25-(OH)₂D₃ (%)</th>
<th>Egg production 4 wk (g)</th>
<th>Egg specific gravity 4 wk</th>
<th>Plasma Ca (µg/dL)</th>
<th>Minerals P (%)</th>
<th>Bone ash (%)</th>
<th>Phytate P retention (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.8</td>
<td>0.33</td>
<td>0</td>
<td>67&lt;sup&gt;abc&lt;/sup&gt;</td>
<td>64.0</td>
<td>1.069</td>
<td>26.7&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.02&lt;sup&gt;c&lt;/sup&gt;</td>
<td>50.1&lt;sup&gt;cd&lt;/sup&gt;</td>
<td>39.6&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>1.8</td>
<td>0.52</td>
<td>0</td>
<td>60&lt;sup&gt;a&lt;/sup&gt;</td>
<td>65.9</td>
<td>1.076</td>
<td>24.6&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>3.05&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>51.2&lt;sup&gt;cd&lt;/sup&gt;</td>
<td>31.3&lt;sup&gt;bc&lt;/sup&gt;</td>
</tr>
<tr>
<td>3.0</td>
<td>0.33</td>
<td>0</td>
<td>72&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>64.7</td>
<td>1.070</td>
<td>22.3&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.91&lt;sup&gt;c&lt;/sup&gt;</td>
<td>49.3&lt;sup&gt;d&lt;/sup&gt;</td>
<td>27.3&lt;sup&gt;ab&lt;/sup&gt;</td>
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<tr>
<td>3.0</td>
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<td>76&lt;sup&gt;b&lt;/sup&gt;</td>
<td>63.2</td>
<td>1.069</td>
<td>31.4&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.44&lt;sup&gt;ab&lt;/sup&gt;</td>
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<td>11.5&lt;sup&gt;b&lt;/sup&gt;</td>
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<td>0.33</td>
<td>5</td>
<td>58&lt;sup&gt;a&lt;/sup&gt;</td>
<td>66.1</td>
<td>1.077</td>
<td>29.5&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>1.76&lt;sup&gt;ab&lt;/sup&gt;</td>
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<td>27.4&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>1.10&lt;sup&gt;ab&lt;/sup&gt;</td>
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</tr>
<tr>
<td>3.0</td>
<td>0.52</td>
<td>5</td>
<td>74&lt;sup&gt;b&lt;/sup&gt;</td>
<td>64.4</td>
<td>0.78</td>
<td>27.8&lt;sup&gt;ab&lt;/sup&gt;</td>
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<td>56.3&lt;sup&gt;a&lt;/sup&gt;</td>
<td>17.8&lt;sup&gt;ab&lt;/sup&gt;</td>
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Mean ± SEM (df = 7): 68 ±7, 64.7 ±1.2, 1.073±0.002, 26.9±2.1, 2.2±0.5, 52.2±0.8, 27.8±8.0, 33.0±8.8

ANOVA (probabilities): df = 0.071

Means with different superscripts are significantly different (p< 0.1).
1,25-(OH)₂ D₃ had any effect on egg production. The week had no effect on egg weight or egg specific gravity. Higher Ca levels in the diet and zero 1,25-(OH)₂ D₃ supplementation both resulted in significantly improved egg weight. The higher Ca level also caused a significant increase in egg specific gravity. A significant interaction between Ca level and 1,25-(OH)₂ D₃ supplementation was obtained. Large increases in specific gravity were obtained when 1,25-(OH)₂ D₃ was fed to the hens receiving the low Ca diet. While the individual levels of Ca and P had no significant effect on the plasma Ca level, there was a significant interaction; raising the P level of the low Ca diet caused a decrease in plasma Ca, while raising the P content of the high Ca diet caused an increase in plasma Ca. Only the P level in the diet caused a significant effect on blood P; low plasma P levels were noted when the low P diet was fed.

All of the individual treatments had a significant effect on bone ash and only one of the interactions (P x 1,25-(OH)₂ D₃) was not significant. The higher bone ash values were obtained from hens fed the high levels of both Ca and P. The greatest effect of 1,25-(OH)₂ D₃ on bone ash was apparent in those hens fed the high Ca and low P diet that was then supplemented with 1,25-(OH)₂ D₃ (49.3% vs 53.9% bone ash).

There is considerable in the phytate P retention data obtained for 14 days and 28 days. In the 14 days data, the most striking effect was the significant decrease in phytate P retention when the high Ca level was present in the diet. While, in the 28 days data, the P level of the diets was the significant factor; at every dietary Ca and 1,25-(OH)₂ D₃ level, increasing dietary P caused a decrease in phytate P utilization.

**DISCUSSION**

There were sufficient significant effects of 1,25-(OH)₂ D₃ supplementation on the diverse criteria measured to indicate definite physiological effects. However, the lack of effect of 1,25-(OH)₂ D₃ supplementation on many criteria, while significant interacts were obtained for the same criteria, indicates the importance of dietary Ca and P levels in any studies of the effects of this compound in laying hens. Harms et al., 1990, conducted several different experiments that explored individually the effect of high and low Ca or P on the response of hens to 1,25-(OH)₂ D₃. The bone ash data obtained by Harms et al., 1990 is similar in most respects to the data obtained in the complete factorial in the paper. When hens were fed a high Ca (3.25%) and low P (0.38%) diet their tibial ash increased from 59.5% to 60.4% and bone breaking strength increased from 4.54 to 6.36 when 1,25-(OH)₂ D₃ was fed. This would be comparable to the results in this trial when 3.0% Ca and 0.33% P were fed and 1,25-(OH)₂ D₃ supplementation resulted in a bone ash increase from 49.3% to 53.9%. Harms et al., (1990), also noted that 1,25-(OH)₂ D₃ supplementation of the high Ca (3.25%) low P diet (0.38%) resulted in a significant decrease in plasma Ca while a slight increase in plasma Ca resulted from 1,25-(OH)₂ D₃ supplementation in the present study. This, of course, could have been caused by small differences in sampling times as related to oviposition (Sooncharernying and Edwards, 1989; Frost et al., 1990) between the different studies. Supplementation of laying diets that are adequate in Ca and P with 1,25-(OH)₂ D₃ both in the present study and in other studies (Harms et al., 1988; Frost et al., 1990; Harms et al., 1990) does not significantly increase tibial ash.

It is interesting that the hens receiving the high Ca diets, had significantly lower 14 days phytate P retention, while those receiving the high P diets had significantly lower 28 days phytate P retention. Concentration of both of these nutrients is known to influence phytate P utilization in broilers (Nelson, 1969; Mitchell and Edwards, 1996). In a similar manner, high dietary Ca and P results in a decrease and increase, respectively, in the retention of phytate P.
Even though the retention of phytate P was not significantly improved by adding 1,25-(OH)$_2$ D$_3$ to the diets of hens receiving the high level of Ca (3.0%) and low P (0.33%), the increase in bone ash was very great. In this case, future studies on the effect of 1,25-(OH)$_2$ D$_3$ on phytate utilization should probably be conducted with the same diet.

REFERENCES


اثر افزودن ۱-۲۵- دی هیدروکسی کوله کلسیفرول در جیره حاوی سطوح مختلف کلسیم و فسفر مرقایی تخم‌گذار

م. شوآزاد. ا. ب. کارلوس. ح. م. ادواردز و م. زاغری

چکیده

اثر افزودن دو سطح (صفر و ۵ میکروگرم در کیلوگرم) ۱-۲۵- دی هیدروکسی کوله کلسیفرول به جیره مرغان تخم‌گذار حاوی سطوح بالا و پایین کلسیم (۲/۳ و ۱/۸٪) و فسفر (۱۵/۲٪ و ۱۳/۳٪) در یک طرح کاملاً تصادفی به روش فاکتوریل ۲×۲×۲ با ۱۴۴ تیمار و ۱۴۴ مرغ تخم‌گذار برای هر تیمار در فضه‌ای انفرادی (جمعه‌ای) مورد مطالعه قرار گرفت. مرغ‌ها جیره‌های آزمایشی را به مدت ۲۸ روز تغذیه نمودند و اطلاعات مربوط به تغییرات وزن بدن، تولید تخم مرغ، مصرف غذا، وزن تخم مرغ و وزن مخصوص تخم مرغ مورد تجزیه و تحلیل آماری قرار گرفت. در پایان آزمایش خونگیری از مرغ‌ها انجام و سطح کلسیم و فسفر بالعصر خون تیمار و پس از کشتن، استخوان ساق یا چپ آنها جهت تعیین خاکستر استخوان ساق جدا گردید. جیره‌های آزمایشی حاوی ذرت و سویا بود و مقدار ۱/۷۵٪ /آکسید کرم به عنوان نشانگر جهت تعیین میزان اپا کلسیم، فسفر و فسفر فیتیک در روزهای ۱۴ و ۲۸ آزمایش به آن افزوده شد. افزودن از ۱-۲۵- دی هیدروکسی کوله کلسیفرول هیچگونه اثر معنی‌داری روی وزن مرغ‌ها، تولید تخم مرغ و مصرف غذا آنها نداشت اما باعث افزایش وزن مخصوص تخم مرغ در مرغ‌های تعیین شده با کلسیم پایین گردید. اثر تیمار روی خاکستر استخوان معنی‌دار بود و بیشترین تأثیر مربوط به افزودن ۱-۲۵- دی هیدروکسی کوله کلسیفرول به جیره حاوی کلسیم و فسفر بالا بود (۳۶/۳٪ در مقایسه جمعه ۳۲/۵٪). در روزهای ۱۴ و ۲۸ بهترین جیره حاوی کلسیم بالا و فسفر بالا اپا فسفر فیتیک در بدن را کاهش دادند.