Dietary calcium to phosphorus ratio affects postprandial phosphorus concentrations in feline plasma

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Abstract

The impact of dietary phosphorus on chronic renal disease in cats, humans and other species is receiving increasing attention. As Ca and P metabolism are linked, the ratio of Ca:P is an important factor for consideration when formulating diets for cats and other animals. Here, we describe a fully randomised crossover study including twenty-four healthy, neutered adult cats, investigating postprandial responses in plasma P, ionised Ca and parathyroid hormone (PTH) following one meal (50% of individual metabolic energy requirement) of each of six experimental diets. Diets were formulated to provide P at either 0·75 or 1·5 g/1000 kcal (4184 kJ) from the soluble phosphorus salt sodium tripolyphosphate (STPP, Na$_5$P$_3$O$_10$), variable levels of organic Ca and P sources, and an intended total Ca:P of about 1:0, 1:5 or 2:0. For each experimental diet, baseline fasted blood samples were collected prior to the meal, and serial blood samples collected hourly for 6 h thereafter. For all diets, a significant increase from baseline was observed at 120 min in plasma PTH (P<0·001). The diet containing the highest STPP inclusion level and lowest Ca:P induced the highest peaks in postprandial plasma P and PTH levels (1·8 mmol/l and 27·2 pg/ml, respectively), and the longest duration of concentrations raised above baseline were observed at 3 h for P and 6 h for PTH. Data indicate that Ca:P modulates postprandial plasma P and PTH. Therefore, when formulating diets containing soluble P salts for cats, increasing the Ca:P ratio should be considered.

Key words: Plasma phosphorus kinetics: Cats: PTH

Chronic kidney disease is a common disease in adult and elderly cats (1). Dietary phosphorus (P) concentrations have received increased focus as a contributing factor in accelerating kidney disease (2,3). Studies in cats investigating postprandial appearance of P in plasma showed that diets incorporating highly soluble forms of P induced dose-dependent increases in P concentration as well as the Ca and P regulatory parathyroid hormone (PTH) (4). At the highest inclusion of highly soluble P tested, there was also a significant reduction in whole blood ionised Ca (iCa) (5). Although the acute exposure studies were not designed to assess the safety of a diet, the observations summarised above are indicative of being aetiological factors in the development of renal changes observed following longer-term dietary exposure by Alexander et al. (2). Studies utilising cats have shown that sustained increases in either plasma P or PTH, respectively, may also have long-term consequences on Ca and P homoeostasis (5,6). The ability to reduce absorption of P from the intestine is therefore desirable in any species from humans to felines (6,7). Two strategies have been employed previously from a dietary perspective: limiting the inclusion level of highly soluble forms of P (8–9) and/or the manipulation of other minerals to modulate the availability of P (10).

The bioavailability of dietary P is not only influenced by the P source and content of the diet but also by the Ca:P ratio, with an increase from around 1:1 to nearly 2:1 reducing P digestibility and urinary P excretion (11–15). Currently, European regulatory body FEDIAF recommends Ca:P between 1:1 and 2:1 for feline adult maintenance diets (16). The mechanism behind the reduction in P bioavailability with increased dietary Ca:P is thought to be due to the formation of insoluble, and hence non-bioavailable, Ca–Mg–P complexes in the intestinal tract when Ca availability is increased (15,17). Similarly, insoluble complex formation is also likely to be the reason that increasing dietary Mg content has also been shown to decrease P absorption and hence urinary P concentration (18). The solubility of the Ca source in the diet is also important in ensuring its availability to form these insoluble complexes and to bind P (14).

Data reported by Coltherd et al. (4) indicated that despite remarkably similar total and added inorganic P inclusions, postprandial plasma P returned to baseline more quickly (within 3 h).
when cats were fed a diet with a Ca:P > 1 compared to Ca:P < 1. This suggested that there might be an effect of Ca:P on the blunting of postprandial plasma P and PTH responses. However, there were other confounding factors within the diets used; therefore, any conclusions were tentative\(^{(4)}\). Thus, the aim of the current study was to systematically investigate the effect of three levels of total dietary Ca:P – low of about 1.0, moderate of about 1.5 and high of about 2.0 – in diets supplemented with a high (1.5 g P/1000 kcal, 4184 kJ) or low (0.75 g P/1000 kcal, 4184 kJ) level of inorganic P from sodium tripolyphosphate (STPP, Na\(_5\)P\(_3\)O\(_{10}\)) on postprandial plasma P, iCa and PTH.

**Materials and methods**

The research protocol was reviewed and approved by the Waltham Animal Welfare and Ethical Review Body and carried out under the authority of the revised Animals (Scientific Procedures) Act 1986 incorporating EU Directive 2010/63. Cats were extensively trained for and habituated to all procedures. A fully randomised six-diet crossover study was performed with twenty-four neutered adult domestic shorthair cats, each cat acting as its’ own control, as previously described\(^{(4)}\). Briefly, the study comprised collection of baseline blood samples from the cats in the fasted state (>18 h), a single 20-min meal exposure to each test diet comprising 50% of the individuals’ daily metabolisable energy requirement, a pattern to which the cats were habituated to, and subsequent hourly blood collection via a cephalic catheter for monitoring of the postprandial plasma levels of P, iCa and PTH for 6 h post-ingestion. Metabolisable energy requirement was calculated as 77·6 × Body Weight\(^{0.731}\) kcal (4184 kJ) per d. The diets offered were blinded to all personnel involved in feeding, sampling the cats and during all sample analysis. Cats were split into six groups of four for sampling. A minimum of 10 d (no more than 14 d) was scheduled between exposures, during which cats were fed a rotation of complete and balanced commercial diets from a range of manufacturers. The rotation coupled with the randomisation of the experimental diets was intended to minimise any increased variability by removing order effects of diets. A minimum of 10 d between samples were implemented for welfare reasons. Our pilot data showed no effects on plasma P, PTH and whole blood iCa of a 7-d break, regardless of diet order, between sampling days\(^{(4)}\).

**Animals**

For details of the cats and diet intakes, see Table 1. To ensure suitability for the study, the cats underwent a pre-study health assessment, including a physical examination by a registered Veterinary Surgeon and haematological, plasma biochemical and urine analysis to confirm the absence of underlying conditions. Cats were group housed in social rooms under routine husbandry conditions. On the day of sampling, cats were allowed to return to their social rooms and monitored closely to ensure the cats’ welfare (i.e. not pulling out the catheter). Throughout the study, commercial rotation (wet and dry format) diets were offered in amounts required to maintain an ideal body weight and body condition score, assessed according to a 9-point scale. These were used to calculate individual metabolisable energy requirement\(^{(19)}\). Deionised water was available ad libitum during the sampling days.

**Diets**

Experimental wet format diets were specifically formulated and manufactured using standard commercial raw materials by Mars Petcare Inc. Poultry necks, pork liver, meat organs, blood plasma, potassium chloride, choline chloride, xylose, xanthan gum, maize starch, and vitamin, amino acid and mineral mixes were the raw materials used for all experimental diets. Poultry necks, pork liver, meat organs, STPP and calcium carbonate were adjusted or added to the formulation to achieve the desired nutritional profiles. Nutritional composition was confirmed through analyses carried out at Eurofins Ltd, Wolverhampton, UK, utilising AOAC official methods of analyses (see Table 2 for details). STPP was used as the main source of inorganic P. Additional inorganic P present in the raw materials accounted for less than 2.0% of the total P from STPP added and was therefore negligible. The diets incorporating high STPP (A–C), contributing 1.5 g P/1000 kcal (4184 kJ), were formulated from the same base recipe, whereby calcium carbonate content was manipulated to achieve the desired total Ca:P ratios of about 1·0, 1·5 and 2·0. The second base diet, in which the levels of the same ingredients used for diets A–C were adjusted, was formulated to contain low STPP (D–F) contributing 0·75 g P/1000 kcal (4184 kJ). Although added P levels were controlled, variability in raw materials resulted in slightly higher total P concentrations in diets E and F, and Ca:P ratios of 0·95, 1·62 and 2·03 for diets A–C and 1·11, 1·70 and 2·04 for diets D–F (Table 2). Diets were randomised to prevent any order bias.

**Blood sampling**

Following an overnight fast (>18 h), topical local anaesthetic (1 ml EMLA™ cream 5 % AstraZeneca UK Limited) was applied and a cephalic catheter positioned, which remained in place for the duration of the sampling period on each sampling day. To maintain patency, a 0·2-ml heparin flush and discard was performed before each collection with a 2-ml saline flush thereafter. A fasted blood sample (1·7 ml) was collected to provide a baseline, with further samples taken at 60 min intervals (0·5 ml at 1, 3 and 5 h and 1·5 ml at 2, 4 and 6 h) following the meal up to and including, 6 h post-feeding.

**Blood sample processing and analysis**

Blood samples (0·5 ml) were collected into lithium heparin tubes (Sarstedt Inc.) and placed on ice until analysis was performed (within 60 min of sampling). iCa was measured in whole blood via an ion-selective electrode using a Stat Profile Prime Veterinary Critical Care Analyser (Nova Biomedicals UK). Performance of the instrument was verified by the analysis of the manufacturers’ tri-level quality control material before and after the sample measurement. Plasma was isolated by centrifugation at 2000 × g for 10 min at 4°C. Phosphorus, in the form of orthophosphate, was photometrically quantified in non-ashed plasma using an AU480 clinical chemistry analyser (Beckman
Postprandial plasma phosphorus kinetics in cats

Table 1. Description of the cats and experimental diet intakes. The values are means and the respective ranges

<table>
<thead>
<tr>
<th>Age (years):</th>
<th>5–14</th>
<th>2.7–7.9</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td>17 males, 7 females</td>
<td></td>
</tr>
<tr>
<td>Body weight (kg)</td>
<td>4.45</td>
<td>3.34–5.62</td>
</tr>
<tr>
<td>Diet intake (g) of each experimental diet (A–F):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A: 148.1 (75–199); B: 134.6 (73–181); C: 139.7 (63–189); D: 140 (73–189); E: 152.8 (98–202) &amp; F: 155.8 (95–207)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>100 % experimental meal* consumed (number of cats)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A: 17; B: 17; C: 16; D: 19; E: 16; F: 18</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Experimental meal = 50 % of the daily metabolisable energy requirement.

Table 2. Nutrient composition of the experimental diets. Units are g/1000 kcal (4184 kJ) metabolisable energy unless otherwise stated

<table>
<thead>
<tr>
<th>Diet</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total P</td>
<td>3.73</td>
<td>3.71</td>
<td>3.84</td>
<td>3.63</td>
<td>4.40</td>
<td>4.58</td>
</tr>
<tr>
<td>Added P†</td>
<td>1.50</td>
<td>1.50</td>
<td>1.50</td>
<td>0.75</td>
<td>0.75</td>
<td>0.75</td>
</tr>
<tr>
<td>Ca</td>
<td>3.55</td>
<td>6.02</td>
<td>7.78</td>
<td>4.03</td>
<td>7.49</td>
<td>9.32</td>
</tr>
<tr>
<td>CaP‡</td>
<td>0.95</td>
<td>1.62</td>
<td>2.03</td>
<td>1.11</td>
<td>1.70</td>
<td>2.04</td>
</tr>
<tr>
<td>Na</td>
<td>2.73</td>
<td>2.51</td>
<td>2.66</td>
<td>1.78</td>
<td>1.94</td>
<td>1.87</td>
</tr>
<tr>
<td>Mg</td>
<td>0.16</td>
<td>0.17</td>
<td>0.18</td>
<td>0.18</td>
<td>0.21</td>
<td>0.22</td>
</tr>
<tr>
<td>K</td>
<td>3.57</td>
<td>3.50</td>
<td>3.68</td>
<td>3.32</td>
<td>3.50</td>
<td>3.25</td>
</tr>
<tr>
<td>Cl</td>
<td>2.87</td>
<td>2.62</td>
<td>2.72</td>
<td>2.60</td>
<td>2.65</td>
<td>3.56</td>
</tr>
<tr>
<td>Vitamin D₃ (IU/1000 kcal [4184 kJ])</td>
<td>452.39</td>
<td>367.16</td>
<td>383.81</td>
<td>318.73</td>
<td>401.17</td>
<td>382.12</td>
</tr>
<tr>
<td>Protein</td>
<td>121.37</td>
<td>128.63</td>
<td>125.78</td>
<td>131.39</td>
<td>136.51</td>
<td>136.88</td>
</tr>
<tr>
<td>Fat</td>
<td>50.57</td>
<td>47.46</td>
<td>46.68</td>
<td>45.53</td>
<td>45.97</td>
<td>48.48</td>
</tr>
<tr>
<td>Ash</td>
<td>25.97</td>
<td>27.47</td>
<td>32.42</td>
<td>22.12</td>
<td>30.64</td>
<td>35.65</td>
</tr>
<tr>
<td>Crude fibre</td>
<td>2.73</td>
<td>3.75</td>
<td>2.59</td>
<td>3.90</td>
<td>6.96</td>
<td>14.26</td>
</tr>
<tr>
<td>Moisture (g/100 g)</td>
<td>83.87</td>
<td>81.90</td>
<td>82.20</td>
<td>82.50</td>
<td>83.20</td>
<td>82.80</td>
</tr>
<tr>
<td>PME§ (kcal/100 g)</td>
<td>73.20</td>
<td>80.10</td>
<td>77.10</td>
<td>76.90</td>
<td>71.80</td>
<td>70.10</td>
</tr>
<tr>
<td>Food base excess (mEq/kg)</td>
<td>29.66</td>
<td>133.22</td>
<td>201.12</td>
<td>18.76</td>
<td>121.53</td>
<td>143.58</td>
</tr>
</tbody>
</table>

IU, international units; PME, predicted metabolisable energy.
- * Suspected analytical error for crude fibre has likely led to an underestimation of PME and an overestimation of g/1000 kcal (4184 kJ) values.
- † Added P represents the target inclusion level of STPP; sodium tripolyphosphate Na₅P₃O₁₀.
- ‡ Ca:P ratio.
- § Calculated by proximate analysis to PME according to NRC 2006(5). 1 kcal = 4.184 kJ.
- † Food base excess calculated from methionine and cysteine according to Jeremias et al. on a per kg food basis(61).

Coulter (UK) Ltd) according to the manufacturer’s instructions, and concentrations reported in mmol/l. To confirm acceptable instrument performance, the manufacturer’s bi-level quality control material was analysed before, during and after the sampling period.

Internal fasted normal ranges for plasma P were 0.9–2.2 mmol/l and for whole blood iCa were 1.19–1.33 mmol/l. Cats are classed as clinically hypercalcaemic when whole blood iCa is above 1.6 g mmol/l(20).

Determination of plasma PTH concentration was performed at the Royal Veterinary College, London (UK). Blood samples (1 ml) were collected in EDTA containing tubes (Sarstedt Inc.), and plasma prepared by centrifugation at 2000 x g for 10 min at 4°C before being stored at −80°C until analysis. Plasma concentrations were measured using a RIA (Skanibodies Laboratory, Inc.) according to the manufacturer recommendations. This assay measures human intact PTH and has previously been validated for use with feline samples, reference range 2.9–26.3 pg/ml(21,22).

Statistical powering

Sample size analyses were performed to determine the number of cats needed in a crossover study design with six diets, to have > 80 % power to detect 20 % change in plasma P levels over seven time points. Twenty-one cats were required, and three extra cats were added to ensure full power was maintained in the event of cats being unable to complete the full six-diet exposure. Plasma P levels from a pilot study(4) were used to estimate the between cat variability and within diet within cat variability. Data were then simulated (1000 for each scenario) according to these variability estimates and by inducing a difference for one diet at one time point. The power analysis was performed, as per the postprandial pilot data statistical analysis (given below). The power was estimated by the percentage of simulations where all the postprandial pilot data statistical analysis (given below). The power was estimated by the percentage of simulations where all comparisons relative to the mean that was changed were significant, at the overall family-wise error rate of 5 %.

Statistical analysis

For each diet, postprandial plasma P (primary measure), iCa and PTH concentrations were analysed using linear mixed effects models, with time nested in diet nested in animal as the random effects, and diet, time point and their interaction as the categorical fixed effects. Plasma P and PTH concentrations were log10 transformed prior to analyses where increasing variation in residuals with increasing value was observed. At some time points,
data were missing due to non-compliant cat behaviour, the catheter becoming unviable or failure of laboratory equipment. However, the restricted maximum likelihood method of analysis was used to account for this and assumed these values were missing at random. Comparisons were made, using a family-wise adjusted 5% level for each measure, between diets at each time point and between baseline to each subsequent postprandial time point for each diet. Comparisons were made within STPP dietary inclusion level (A v. B v. C and D v. E v. F) and within Ca:P level (A v. D, B v. E, C v. F). Accordingly, means, difference between means (or fold change on log10 transformed data) were reported as differences with 95% CI and are reported for all diets. Comparisons between diets are observed in cats fed diets B and C was not found (P > 0.05). Significant reductions in plasma P concentrations from baseline were observed in cats fed diets E and/or F postprandially, with the exception of 120 min, which did not differ from baseline (P > 0.05). Reductions from baseline values were also observed in cats fed diets B and C at 300 and 360 min postprandially (P < 0.05).

When the AUC (Fig. 2) for each response profile in postprandial plasma P (Fig. 1) were determined, highest values were observed when cats were fed diet A (P < 0.001) compared with all other relevant diet exposures. Within each subset of diets, greater AUC values were observed when cats were fed diets A compared with B and C, and diet D compared with E and F, indicating an inverse relationship to Ca:P ratio level (P < 0.001 for all). However, no significant differences were observed when cats were fed the diets containing the moderate compared with high Ca:P ratio (diets B v. C and E v. F, P > 0.05). Across the diet subsets, AUC values were greater when cats were fed diets A v. D, B v. E and C v. F (P < 0.001 for all) suggesting a dose-dependent response to dietary STPP inclusion level. The mean AUC calculated for the postprandial plasma P responses to diets E and F were negative (Fig. 2).

Effect of dietary inorganic phosphorus and calcium to phosphorus ratio on postprandial whole blood ionised calcium

Postprandial blood iCa concentrations (Fig. 3) significantly decreased from baseline in all cats following intake of each diet (P < 0.001). Diets A, D and E were also significantly lower than baseline concentrations at the final time point (P < 0.001, P = 0.003 and P = 0.001, respectively). At 240 min post-meal, a significantly lower iCa level was observed in cats after the exposure to diet A compared with diets C and D (P < 0.001). Among diets with high STPP supplementation, significant decreases were observed at 240 and 300 min post-meal when cats were fed diet A with the low Ca:P ratio compared with diet C with the high Ca:P ratio (P < 0.001). Within the subset of diets with low STPP supplementation, an effect of Ca:P level was not observed (P > 0.05).

When the AUC (Fig. 4) for each response profile in postprandial iCa (Fig. 3) were determined, the values calculated were negative due to the decreased postprandial iCa levels after the exposure to each diet compared with baseline. Diet A induced the largest AUC (A v. B P = 0.025, A v. C P < 0.001 and A v. D P < 0.001, Fig. 4), while diet B had a significantly larger AUC than diet E (P = 0.027).

Effect of dietary inorganic phosphorus inclusion and calcium to phosphorus ratio on postprandial plasma phosphorus

Postprandial responses in plasma P were increased, compared with baseline (fasted state), from 120 min post-feeding experimental diets A-D to cats (P < 0.001; Fig. 1). These increases in postprandial plasma P reached the maxima from baseline at 180 min in the cats fed the diet subsets containing low Ca:P (A and D) and returned to baseline at 300 min. Cats fed the mid to high (B and C, respectively) dietary Ca:P had a maxima postprandial plasma P concentration at 120 min which returned to baseline at 240 min (P < 0.001; Fig. 1). Peak plasma P concentrations of 1.80, 1.53 and 1.50 mmol/l were observed for diets A, B and C, respectively, compared with 1.54, 1.28 and 1.16 mmol/l for diets D, E and F, respectively. A significant difference between the plasma P-values from 120 min onwards when fed diets B and C was not found (P > 0.05). Significant reductions in plasma P concentrations from baseline were observed in cats fed diets E and/or F postprandially, with the exception of 120 min, which did not differ from baseline (P > 0.05). Reductions from baseline values were also observed in cats fed diets B and C at 300 and 360 min postprandially (P < 0.05).
Fig. 1. Mean plasma phosphorus (mmol/l) concentrations over 360 min postprandial periods in cats following a meal of each experimental diet A–F comprising 50 % of the daily metabolisable energy requirement. *Significant differences to the baseline fasted sample (time 0); \( P \leq 0.05 \). The vertical bars for each mean indicates 95 % CI. STPP, sodium tripolyphosphate.

Fig. 2. Mean calculated AUC (h.mmol.l\(^{-1}\)) values for postprandial plasma P responses in cats over 360 min periods following intake of a meal for each experimental diet A–F comprising 50 % of daily metabolisable energy requirement (MER) *A significant difference between diet exposure responses; \( P \leq 0.05 \). The vertical bars for each mean indicates 95 % CI. STPP, sodium tripolyphosphate.
Fig. 3. Mean whole blood ionised calcium (mmol/l) concentrations over 360 min postprandial periods following intake of a meal for each experimental diet A–F comprising 50 % of the daily metabolisable energy requirement (MER). *Significant differences to the baseline fasted sample (time 0); $P \leq 0.05$. The vertical bars for each mean indicates 95 % CI. STPP, sodium tripolyphosphate.

Fig. 4. Mean calculated AUC (h.mmol.l$^{-1}$) values for postprandial whole blood ionised calcium responses in cats over 360 min periods following intake of a meal for each experimental diet A–F comprising 50 % of the daily metabolisable energy requirement (MER). *Significant difference between diet exposure responses; $P \leq 0.05$. The vertical bars for each mean indicates 95 % CI. STPP, sodium tripolyphosphate.
Effect of dietary inorganic phosphorus and calcium to phosphorus ratio on postprandial plasma parathyroid hormone

Compared with baseline, all diets produced an increase in postprandial plasma PTH at 120 min ($P < 0.05$, Fig. 5). The maxima in postprandial plasma PTH responses in cats (Fig. 5) differed, with the highest peak observed at 240 min after diet A intake compared with the other diets’ lower peaks reached at 120 min ($P < 0.001$ for each comparison). At the lower STPP inclusion level, diet D exposure induced significantly higher values at 240 min compared with diets E and F with medium and high Ca:P, respectively ($P < 0.001$). The plasma PTH responses to diet A and diet D were still raised above baseline levels at 360 min following the meal ($P < 0.001$ and $P = 0.008$, respectively).

Within diet subsets of STPP supplementation, AUC values calculated for each response profile in postprandial PTH (Fig. 6) also indicated significant increases after diet A intake compared with diets B and C ($P < 0.001$), as well as between D and F ($P < 0.001$). There was no significant difference between diet D and E ($P = 0.083$). Across diet subsets, dose-dependent responses to dietary STPP inclusion level were observed with significantly higher values observed following intake of diet A compared with D ($P < 0.001$) with low Ca:P, diet B compared with E ($P = 0.01$) with moderate Ca:P and diet C compared with F ($P < 0.001$) with high Ca:P.

Discussion

To better understand the role of dietary total Ca:P in P metabolism and regulatory (PTH) responses in the healthy, adult cat, a single-meal postprandial absorption kinetics model, as indicated by blood and plasma responses, was utilised. The data indicate that increasing dietary Ca:P from about 1:0 to between 1:6 and 2:2, reduced the peak and duration of both plasma P and PTH elevation in both subsets of diets containing two levels of STPP supplementation. The data support our previous observation that Ca:P ratio acutely affects P appearance in the plasma (14), reflecting the gross changes observed in both apparent digestibility and urinary P excretion in other studies (11-15). This finding adds further credence to the hypothesis that lower dietary Ca:P may be an additional factor in the development of reduced renal function and renal tissue changes reported in two long-term studies investigating the effects of dietary P levels in cats (2). Furthermore, the level of soluble P salt supplementation (STPP) also significantly and dose-dependently affected postprandial P, iCa and PTH responses, as previously reported (4).

In other animal species, low dietary Ca:P ratios have been associated with changes in Ca and P absorption, metabolism and regulation, as well as associated renal function and structure changes. Hock et al. (27) found that increasing total dietary Ca from 0.5-2% weight/weight, wt/wt and decreasing Ca:P to 0.6 increased severity of nephrocalcinosis in rats when compared with a diet with a Ca:P of 1:3 and a total Ca inclusion of 25% wt/wt. The rats with the most severe nephrocalcinosis also had higher urinary P excretion, whilst urinary Ca excretion was relatively stable and instead was eliminated from the body via the faeces (27). This elimination of increased dietary Ca in the faeces has also been observed for cats (28).

In cats experiencing renal secondary hyperparathyroidism, those assigned a dietary regimen involving phosphate restriction and a higher Ca:P (1:65 and 1:89, respectively) had reduced PTH and plasma P compared with cats assigned a maintenance diet (Ca:P = 1:11) (29). Paflack et al. (30) increased dietary bone meal inclusion to produce diets with a Ca:P of 0.76:1, 1:05:1 and 1:28:1, also resulting in increasing total Ca and P concentrations, in order to assess formation of uroliths in cats. Data showed that faecal elimination of P was increased when Ca:P was 1:28:1, whilst urinary excretion (% excretion by intake) and apparent digestibility of P were lowest on this
diet(30). Data also showed that urinary Ca excretion (% excretion by intake) and PTH concentrations were significantly raised when the cats were fed the diet with Ca:P of 0.76:1(30).

In a human study, Calvo et al. (31) offered participants a test diet regime with a mean Ca:P of 0.25 compared with a control regime with a Ca:P of 1.0 in the four meals eaten over the course of a day. This resulted in postprandial P, iCa and PTH kinetics comparable to the data presented herein(31). However, as this appears to be a cumulative effect following multiple meals, it is difficult to compare directly with our results following one meal.

For whole blood iCa, diet A caused the lowest iCa concentration at 240 and 300 min and highest PTH values at 240 min. These data suggest that the reduction in whole blood iCa after the consumption of diet A may be associated with a homeostatic up-regulation of PTH to a higher degree than for the other experimental diets in order to return iCa concentrations to baseline(32). Similar findings were observed previously(4), where the experimental diet associated with the lowest postprandial iCa and highest PTH also had a low Ca:P (0.6) and 3.5 g/1000 kcal (4184 kJ) of highly soluble P in the form of sodium dihydrogen phosphate (SDHP, NaH2PO4). This may indicate that low dietary Ca:P (<1) and a concomitant high inclusion of inorganic P (≥1.5 g/1000 kcal(4184 kJ)) causes a larger response in PTH to regulate iCa concentrations. This may help explain the adverse effects to renal health seen in cats following 4 weeks of exposure to such a diet in Alexander et al.(2).

Decreased iCa and/or increased serum P over prolonged periods have been linked to increased PTH concentrations and parathyroid dysfunction in humans(33). It should be noted that the dietary Ca of the experimental diets did differ, with diets A–C having a lower content than their respective counterparts (D–F). A study in humans found that a high Ca, high vitamin D breakfast increased iCa postprandially compared with a low Ca, low vitamin D breakfast(50). This study did not find any significant difference in PTH postprandially between the diets(50).

Significantly higher, dose-dependent changes in plasma P and PTH concentrations were observed after the intake of diets with increasing STPP inclusion in this and previous studies(4). The data from the current study also indicate an interactive effect of STPP inclusion level with total dietary Ca:P ratios, with increasing Ca:P to moderate (about 1.5) or high (about 2) values apparently leading to decreased appearance of P in the plasma following a meal at both STPP inclusion levels when compared with low dietary Ca:P (about 1). The magnitude of peak postprandial plasma P concentrations following diet D intake (low STPP, low Ca:P) was similar to those after the intake of diets B and C (high STPP and moderate and high Ca:P, respectively), resulting in similar AUC. However, the maximum value reached following intake of both diets A and D, with Ca:P ratios of about 1 but differing in STPP inclusion, was at 180 min v. 120 min following intake of all the other diets with higher Ca:P ratios. Thus, these data indicate that the observed effect of diets including a soluble P salt in the form of STPP, which raised plasma P and PTH concentrations, could be moderated by increasing the dietary Ca:P between about 1.5 and 2.0. This could mean that adjustments in total Ca:P ratio to ≥1:6 in dietary formulations

Fig 6. Mean calculated AUC (h.mmol.l⁻¹) values for postprandial plasma parathyroid hormone responses over 360 min periods following intake of a meal for each experimental diet A–F comprising 50% of the daily metabolisable energy requirement (MER). *Significant difference between diet exposure responses; P ≤ 0.05. The vertical bars for each mean indicate 95% CI. STPP, sodium tripolyphosphate, PTH, parathyroid hormone.
may aid in ameliorating increases in plasma P and PTH. This hypothesis is supported by the results from a recent 7-month feeding trial in cats testing STPP inclusion levels providing an inorganic P level of 1·0 g/1000 kcal (4184 kJ), but with two different total P and Ca:P ratios\(^{35}\). Although a no-observed-adverse-effect level was observed, cats fed the diet with the lower total P and Ca:P ratio showed some increases in measures related to Ca and P metabolism, namely urinary excretion of P and plasma FGF-23 concentration, that were not found in cats fed the diet with the higher total P and higher Ca:P ratio\(^{35}\).

When comparing diets with similar formulations across this study with previous acute\(^{40}\) and chronic exposure\(^{22}\) studies, some differences in responses were observed. Despite similar dietary total and soluble P levels and Ca:P ratios, results reported in our previous study\(^{40}\) showed a lower postprandial maximum of 1·57 mmol/l in plasma P that did not return to baseline by 360 min post-feeding. In addition, no postprandial PTH peak was observed with levels not differing from baseline within the 360-min postprandial period. And yet when that diet\(^{40}\) was fed to cats for 28 weeks, development of renoliths and structural changes in the kidneys were observed in a number of cats\(^{2,4}\). The formulation of the two diets in the current compared with previous studies\(^{2,3,4}\) differed slightly, however, and this suggests that there are several other diet-related factors that could affect the postprandial levels of P and PTH in the plasma of cats following a meal. These factors could include differences in the production technology and diet format, the raw materials and ingredients used, fibre content, other minerals, base excess and vitamin D concentration. Data from cats, fed various cellulose fibre lengths, showed that increased dietary crude fibre content may increase Ca, P and Mg excretion, leading to lower absorption\(^{36}\). Dietary fibre may also change the pH of the ileal digesta, increasing Ca absorption thus possibly leading to a reduced availability to form insoluble Ca–Mg–P complexes in the intestine, thereby indirectly affecting P absorption\(^{15}\).

Commercially relevant diet formulations rely on varying combinations of raw materials to reach desired nutrient compositions. These raw materials can vary in nutrient profile from batch to batch\(^{37}\) and therefore cause unintended differences between the experimental diets. However, the current study used wet format diets, and raw materials are unlikely to contain significant phytate or fibre sources. If the raw materials did contain these fibre sources, diets A–C should have followed the same pattern as diets D–F; therefore, the unexpectedly high crude fibre content of F cannot be accounted for. A point to note is that any potential analytical error in crude fibre would also lead to an underestimate of the PME of the diet, leading to (1) an overestimation of P in g/1000 kcal (4184 kJ) for diet F, and (2) overfeeding the cats diet F relative to the diets A–E. These factors should be taken into account when reviewing the data. Despite this uncertainty, the findings are more relevant to commercial formulations than employing semi-purified diet formats where nutrient digestibility is significantly greater. The diets formulated for this study increased in base excess with increasing dietary Ca:P, which limits our ability to separate out the two factors and should therefore be a consideration for future studies. Vitamin D is linked to increased P absorption in cats\(^{38}\) and, although attempts to maintain concentrations were successful for the most part, the higher vitamin D content in the current diet A may have led to the higher peak in plasma P compared with the more prolonged but smaller postprandial increases in plasma P observed previously\(^{41}\). Diet format may also alter the absorption rate of minerals and affect plasma P postprandial kinetic profiles, likely linked to differences in raw ingredients and often higher fibre content in dry v. wet diets as described above. Previous investigation of the postprandial plasma P kinetics of two dry format diets, one containing SDHP and the other STPP, showed no significant differences in peak height and duration of plasma P; however, the AUC was significantly higher in the SDHP diet\(^{41}\). SDHP has a higher water solubility than STPP (800 g/l and 150 g/l, respectively)\(^{39}\) most likely contributing to the AUC data differences. Renal P excretion has also been found to be increased in cats fed a diet with sodium monophosphate compared to cats fed a diet containing calcium monophosphate\(^{10}\). These data may suggest that there are differences in absorption between added phosphorus salts that warrants further attention. The use of a wet and dry commercial diet rotation, which is part of the cats’ normal care routine, between experimental trials may have influenced the outcome of the trial due to the potential for exposure to different dietary vitamin D and mineral contents, possibly altering the homeostatic capacity of PTH\(^{31,40}\). There were no observed differences in baseline (fasted) values of plasma P, PTH or whole blood iCa between exposures to the experimental diets. This suggests that the randomisation of the experimental diets fed coupled with the 18-h fast and the crossover design may have limited variation between fasted values. However, a potential impact of the dietary regimen between sampling periods cannot be completely discounted.

While there are still some unknowns around the possible contributions of other dietary factors, such as added soluble P source, our previously reported data indicate a contributory role of a low Ca:P of less than 1·0 in the adverse effects reported when single-batch experimental diets were fed to healthy adult cats for a minimum of 4 weeks\(^{2}\). Our previous data indicate that this effect may already be seen at a ratio of 1·5. However, identifying the role of dietary Ca:P was not the goal of those studies\(^{4}\). The preliminary findings here suggest that the effects of dietary soluble P source STPP on plasma P and PTH concentrations can be mitigated by increasing the Ca:P ratio to 1·6 and above, as well as by reducing the level of added soluble P in the diet. These factors should be a consideration for formulation of feline diets and would further benefit from longer-term feeding trials, including measures of renal function, to verify.

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Supplementary material

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References


