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PHOSPHORUS DIGESTIBILITY

Results of an international phosphorus digestibility ring test with broiler chickens

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1 **ABSTRACT** The objective of this ring test was to investigate the prececal phosphorus (P)
2 digestibility of soybean meal (SBM) in broiler chickens using the trial protocol proposed by
3 the World's Poultry Science Association. It was hypothesized that prececal P digestibility of
4 SBM determined in the collaborating stations is similar. Three diets with different inclusion
5 levels of SBM were mixed in a feed mill specialized in experimental diets, and transported to
6 17 collaborating stations. Broiler chicks were raised on commercial starter diets according to
7 station-specific management routine. Then they were fed the experimental diets for a
8 minimum of 5 d before content of the posterior half of the ileum was collected. A minimum
9 of 6 experimental replicates per diet was used in each station. All diets and digesta samples
10 were analyzed in the same laboratory. Diet, station, and their interaction significantly affected
11 ($P < 0.05$) the prececal digestibility values of P and calcium of the diets. The prececal P
12 digestibility of SBM was determined by linear regression and varied between stations from 19
13 to 51 %, with significant differences among stations. In a subset of 4 stations, the prececal
14 disappearance of *myo*-inositol 1,2,3,4,5,6-hexakis (dihydrogen phosphate)-P; InsP₆-P) was
15 also studied. The prececal InsP₆-P disappearance correlated well with the prececal P
16 digestibility. We hypothesized that factors influencing InsP₆ hydrolysis were main
17 contributors to the variation in prececal P digestibility between stations. These factors were
18 probably related to the feeding and housing conditions (floor pens or cages) of the birds in the
19 pre-experimental phase. Therefore, we suggest that the World's Poultry Science Association
20 protocol for the determination of digestible P be extended to the standardization of the pre-
21 experimental period. We also suggest that comparisons of P digestibility measurements
22 between studies are only made with great caution until the protocol is more refined.

23 **Key words:** broiler chickens, phosphorus, phytate, prececal digestibility, variability

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INTRODUCTION

Phosphorus (P) is an element of high relevance for poultry feeding. Diets are usually supplemented with feed phosphates or phytase or both in order to fulfil the animal's requirement for available P. However, excessive intake can contribute to environmental problems in areas with high livestock density. The variation in P availability of different feed raw materials is high (Shastak and Rodehutsord, 2015), and it is generally accepted that the use of P as a globally finite resource can be optimized by considering the differences that exist in P availability of feed raw materials.

Different response criteria and descriptive terms for available P have been used in the literature of the past seven decades (Shastak and Rodehutsord, 2013). These differences make it difficult to compare results obtained by using different techniques in different laboratories, and to compile comprehensive feedstuff tables needed by the industry. In an attempt to improve this situation the Working Group No 2: Nutrition of the European Federation of Branches of the World's Poultry Science Association proposed a standard protocol for the determination of P availability (WPSA, 2013). This protocol is based on using the digestibility measured at the terminal ileum of broiler chickens (prececal digestibility of P (**pcdP**), otherwise also referred to as ileal digestibility). The protocol defines assay details relevant for the outcome of the measurement, such as age of birds, minimum number of experimental replicates, diet composition, and P and calcium (Ca) levels in the diet.

The expectation from implementing this standard protocol is that results generated in different research stations for the same feed raw material are similar and thus can better be compared and used by the industry. However, application of the standard protocol has not been compared yet between stations. Thus, our objective was to compare the results between our laboratories for one feed raw material when based on the WPSA (2013) standard protocol.

49 We choose soybean meal (**SBM**) for this comparison, and we hypothesized that the pcdP of
50 SBM determined in our laboratories is similar.

51

52 **MATERIALS AND METHODS**

53 Seventeen research stations from Europe and North America collaborated in this study and
54 determined the pcdP of SBM following the principles of the standard protocol of WPSA
55 (2013), which includes regression analysis. The concept of this ring test required that all
56 stations use the same diets and thus variability in raw material quality and diet preparation
57 was mitigated. Furthermore, chemical analyses of all diets and ileal digesta collected in all
58 stations were conducted in the same laboratory. Customs regulations related to cross-
59 continental shipment of feed samples made it impossible for more laboratories from other
60 regions of the world to participate in the ring test.

61

62 ***Experimental diets***

63 Three diets that differed in the inclusion level of SBM (diets A, B, C) were formulated based
64 on the examples of WPSA (2013) (Table 1). The de-hulled, solvent-extracted SBM was
65 included at the expense of corn starch, thus making SBM the only source of variation in P
66 content of the diets. Analytical characterization of the SBM is presented as a footnote in Table
67 1. Titanium dioxide was included (0.5 %) as the indigestible marker. The concentrations of
68 total P and phytic acid (*myo*-inositol 1,2,3,4,5,6-hexakis (dihydrogen phosphate)-P; **InsP₆-P**)
69 in the diets ranged from 3.0 to 4.6 and 1.6 to 2.3 g/kg dry matter (DM), respectively. The
70 Ca:total P ratio was analyzed in the diets to be 1.5:1.0, which was very close to the ratio
71 recommended by WPSA (2013) (1.3:1.0 - 1.4:1.0). Diets were manufactured at Research Diet
72 Services (location Wijk bij Duurstede, Netherlands). First, all ingredients except the variable
73 ones (SBM, corn starch and limestone) were mixed in one lot. Then, diets A and C were
74 mixed by adding the respective amounts of SBM, corn starch and limestone. Finally, diet B

75 was mixed by 50 % diet A and 50 % diet C. All diets were pelleted (die hole: 3.0 mm x 17
76 mm) with a little steam addition. Temperature of the pellets measured directly after leaving
77 the die was 77°C for diets A and B and 74°C for diet C. The experimental diets were bagged,
78 sealed, and transported to the 17 participating stations.

79

80 ***Birds and experimental procedures***

81 Animal trials at the participating stations were conducted between March and September
82 2014. All animal procedures were in accordance with the animal welfare regulations that were
83 applicable for the individual participating station. While basic principles of the WPSA (2013)
84 protocol were applied in all laboratories, details of the trials were specific for each station
85 (Table 2). Broiler chicks (Ross 308, Ross 708, Ross PM3, and Heritage 5632) were fed
86 commercial starter feed until they were between 12 and 23 d old and when the treatment diets
87 were introduced. At least 6 experimental replicates were used per diet by each station, and
88 each replicate had a minimum of 8 birds. One laboratory used only 4 birds per replicate but 10
89 replicates per diet. Feed and drinking water were offered for ad libitum consumption. The
90 ADFI and average BW were recorded. Birds were fed the experimental diets for a minimum
91 period of 5 d and then were sacrificed at an age of 21-28 d. The abdominal cavity was
92 immediately opened, the digestive tract removed, and the ileum (section between Meckel's
93 diverticulum and 2 cm anterior to the ileo-ceco-colonic junction) dissected. The digesta of the
94 distal half of the ileum was obtained by flushing with water or by gentle squeezing. Digesta
95 from all birds of one replicate were pooled into one sample. Samples were dried (freeze-
96 drying or oven drying), ground, and sent to the Hohenheim laboratory together with the
97 respective diet samples for chemical analyses.

98

99 ***Chemical Analyses***

100 Proximate nutrients in the SBM and CP concentration in the diets were analyzed according to
 101 the official methods (Verband Deutscher Landwirtschaftlicher Untersuchungs- und
 102 Forschungsanstalten (VDLUFA, 2007)). Analysis of Ca, P, and Ti in diets and digesta
 103 samples was performed using an inductively coupled plasma optical emission spectrometer
 104 following a sulfuric and nitric acid wet digestion with all specifications described by Zeller et
 105 al. (2015b). Concentrations of InsP₆ and lower inositol phosphates in the diets were analyzed
 106 following EDTA extraction at pH 10 using high-performance ion chromatography as
 107 described by Zeller et al. (2015c). Digesta samples from 4 stations that had freeze-dried the
 108 samples and were found to be different in pcdP values were also analyzed for InsP₆ using the
 109 same method as used for the diets. This was not planned from the beginning and is not part of
 110 the WPSA (2013) protocol. Hence, some stations oven-dried the samples. Oven dried samples
 111 were not considered for InsP₆ analysis because some InsP₆ might be hydrolyzed after
 112 sampling from the ileum during the drying process.

113

114 *Calculations and Statistics*

115 Prececal digestibility of P and Ca, and disappearance of InsP₆ (y) was calculated on a pen
 116 basis according to the following equation:

$$117 \quad y (\%) = 100 - 100 \times \left(\frac{\text{Ti in the diet (g/kg DM)}}{\text{Ti in the digesta (g/kg DM)}} \right) \times \left(\frac{\text{InsP}_6 \text{ or P or Ca in the digesta (g/kg DM)}}{\text{InsP}_6 \text{ or P or Ca in the diet (g/kg DM)}} \right)$$

118 The amount of pcdP (g/kg DM) was calculated by multiplication of the P concentration of the
 119 diet (g/kg DM) and the respective digestibility (%) divided by 100.

120 Statistical evaluation of the data used the MIXED procedure of SAS for Windows
 121 (Version 9.3, SAS Institute, Cary, NC). Evaluation of P digestibility, Ca digestibility, and
 122 InsP₆ disappearance values of the diets was performed using diet (A, B, C), stations (1 - 16),
 123 and their interaction as fixed effects. The trial in one station generated questionable
 124 digestibility data for P and Ca that could not be explained. Hence data of this trial were not
 125 included in the data evaluation and only 16 trials were used.

126 Digestibility of P from SBM was calculated for each of the 16 stations by linear
127 regression. Linear regressions of the type $y = a + mx$ were calculated using the SOLUTION
128 statement to describe the relationship between pcdP content and P content in the diet (both in
129 g/kg DM) for each station. Because differences in the P content between diets originated only
130 from SBM inclusion, calculated slopes multiplied by 100 are regarded as an estimate of the
131 pcdP of SBM (WPSA, 2013). The R^2 and root MS error are reported as measures for the
132 goodness of fit. Differences in pcdP of SMB between stations were compared using the
133 ESTIMATE statement.

134
135

RESULTS

136 The average pcdP of the diets across all stations for diets A, B, and C were 67, 61, and 55 %,
137 respectively (Figure 1). Effects of diet, station, and their interaction were statistically
138 significant ($P < 0.05$). The ranges of pcdP values of the diets across stations were 55 to 82 %
139 for diet A, 46 to 79 % for diet B, and 45 to 69 % for diet C (Table 3).

140 When the pcdP concentration of the diets was regressed against total P concentration,
141 estimated slopes of the linear regressions ranged between stations from 0.19 to 0.51 (Table 4).
142 Even when the two lowest and two highest slopes were disregarded, the slopes of the
143 remaining stations still ranged from 0.22 to 0.42 with significant differences among stations.

144 The average prececal (pc) disappearance of InsP_6 as studied in 4 stations varied from
145 58 to 74 %, 43 to 67 %, and 23 to 58 % for diets A, B, and C, respectively (Table 5), thus
146 confirming the trends in differences seen in pcdP of the diets. Increments in dietary InsP_6 with
147 increasing SBM level of the diet were not related to an increase of pc InsP_6 disappearance
148 (Figure 2). However, across all diets and the 4 stations, a relationship between the pc
149 disappearance of InsP_6 -P and pcdP content became apparent, indicating that a very high
150 proportion of the InsP_6 -P that disappeared until the end of the ileum was absorbed (Figure 3).

151 The average prececal digestibility of Ca (pcdCa) of the diets across all stations was 57
152 % (diet A), 51 % (diet B), and 46 % (diet C), with a similar range between stations as found
153 for pcdP (Figure 1). Effects of diet, station, and their interaction on pcdCa were statistically
154 significant ($P < 0.05$). Within the range studied the relationship between pcdP and pcdCa in
155 the diets was linear and not significantly different between diets (Figure 4). On average, the
156 concentration of pcdCa in the diets increased by 0.97 g/kg DM with each 1.00 g/kg DM
157 increase in pcdP.

158

159 DISCUSSION

160 The underlying hypothesis of this study was that digestibility values determined for the diets
161 and for the SBM are similar among stations. This hypothesis must be rejected on the basis of
162 the results. The pcdP for the same SBM ranged between 19 and 51 % among stations, which
163 can be a frustrating fact from the viewpoint of feed producers aiming to use P digestibility
164 data in their feed formulation matrix. It is of note that the opposite was found in a ring test on
165 pcd of amino acids in broilers (Ravindran et al., submitted). There, pcd of amino acids of a
166 corn-SBM-based diet was similar among the 5 collaborative stations, once important protocol
167 details had been agreed on. Probably there is something specific on the targeted nutrient
168 (amino acids vs. P and InsP_6) that caused high variation in one ring test but not in the other. It
169 cannot be ruled out that this difference is related to hormonal control of absorption known for
170 Ca and P, but not for amino acids. However, for reasons subsequently mentioned other factors
171 are seen more likely involved.

172 In the current ring test we used the same diets in all stations to mitigate variation
173 potentially caused by raw material origin or diet processing. All diet and digesta samples were
174 analyzed in the same laboratory, hence minimizing variation of results potentially caused by
175 differences in analytical protocols. In spite of this standardization, the large differences have
176 occurred. This variation is most likely associated with how the experiment was conducted and

177 birds managed in each station. While most assay details are standardized in the applied
178 protocol (WPSA, 2013), some were not and will be discussed herein.

179 The starter diets used in the pre-experimental period met the bird's requirements, but
180 were specific for each station and not of the same ingredient composition. Digestive capacity
181 adaptation to Ca and P deficiencies or imbalances can occur within 48 h (Angel et al., 2013).
182 Hence, any adaptation to the experimental diets would have already occurred in the present
183 study before samples were taken. However, it cannot be ruled out that starter diet details have
184 affected the measurements subsequently made with the experimental diets. The P
185 concentration of the starter diets (formulated values) was between 0.5 and 0.7 %, and the Ca
186 concentration between 0.7 and 1.1 %. Some of the starter diets contained an added phytase
187 while others did not. Compensatory adaptation in growth and bone mineralization in a later
188 growth phase occurred when broilers were fed a diet moderately deficient in P and Ca from
189 hatching to 18 d (Yan et al., 2005). A recent study showed that a low level of P in the diet fed
190 from 10 to 21 d can affect the mRNA levels of several genes encoding Ca and P transporters
191 at 35 d of age, depending on the amount of P and Ca fed from 22 to 35 d (Rousseau et al.,
192 2016). These authors concluded that chickens are able to adapt to early dietary changes in P
193 and Ca through improvement of digestive efficiency in a later phase. Starter diet P and Ca
194 levels were different across participating stations in the current study, but none was deficient.
195 It cannot be ruled out but not substantiated with the data that the differences affected the
196 results. A negative trend might be interpreted when the P digestibility data are related to the P
197 level of the starter diets (Figure 5A); however, the regression line did not significantly deviate
198 from zero ($P = 0.08$). A relationship with the formulated phytase content of the starter diet
199 also was not apparent (Figure 5B). Because only the formulated values of P, Ca, and phytase
200 are available for the starter diets, and because we do not have information about raw material
201 composition of the starter diets, relationships between starter diet characteristics and
202 determined digestibility values should be viewed with caution. In such complex design,

203 colinearity also between other factors of variation might occur (starter diet, duration of period,
204 age, etc.) and thus could bias interpretation.

205 Some starter diets contained a coccidiostat, while others did not. As a consequence,
206 microbiota colonization of the digestive tract might have developed differently between
207 stations, which could affect InsP₆ hydrolysis as seen in the differences among stations and
208 diets (Table 5). Previous studies have found a relatively high level of prececal or excreta InsP₆
209 disappearance in birds (25 to 78 %) even when diets devoid of any supplemental or plant-
210 intrinsic phytase were used (Amerah et al., 2014; Applegate et al., 2003; Delezie et al., 2012;
211 Li et al., 2016; Tamim et al., 2004; Zeller et al., 2015a; Zeller et al., 2015b). This range of
212 values is high and probably related to dietary Ca levels or level and origin of InsP₆. However,
213 the generally high InsP₆ disappearance values strongly indicate that enzymes of non-feed
214 origin, namely endogenous mucosal and bacterial phytases, have caused a substantial InsP₆
215 hydrolysis. Little is known so far about the contribution of specific bacteria to phytase activity
216 in the digestive tract of chickens. However, lactobacilli were shown to be the main colonizers
217 in the crop, jejunum, and ileum of broilers (Witzig et al., 2015). Some of the lactobacilli
218 strains that were detected in gut content of broilers have been characterized to express high
219 phytase activity (Palacios et al., 2008; Taheri et al., 2009). Use of the coccidiostat monensin
220 reduced abundance of *Lactobacillus* sp. in ceca content (Danzeisen et al., 2011) and nisin-
221 sensitive *L. reuteri* isolates were detected in the crop (Abbas Hilmi et al., 2007). Hence it is
222 possible that coccidiostats included in some but not all starter diets used in the current ring
223 test reduced abundance of bacteria known to possess phytase activity and thus InsP₆
224 hydrolysis. Initial microbiota colonization in the starter phase probably was effective in the
225 experimental phase and contributed to InsP₆ hydrolysis from SBM to a different extent. We
226 observed a very close relationship between InsP₆-P disappearance and pcdP content of the
227 diets (Figure 3), and although P of non-InsP₆ origin in SBM also contributed to the change in

228 pcdP content, the relationship well demonstrated that InsP₆ hydrolysis was a major
229 determinant for the pcdP of SBM.

230 Another factor of potential relevance for microbial colonization and development of
231 the digestive tract is the way birds were kept in the pre-experimental period. When birds were
232 raised in floor pens, litter material intake could affect digestive tract development in general
233 and microbial colonization in particular. Floor pen raising was practiced in some of the
234 stations because required as per approval from the ethics committee or for other reasons.

235 Endogenous enzymes involved in intestinal InsP₆ hydrolysis can also originate from
236 epithelial secretion because studies using purified brush border membrane vesicles from
237 different sections of the small intestine of broiler chickens reported phytase activity (Huber et
238 al., 2015; Maenz and Classen, 1998; Onyango et al., 2006). Significant differences in the V_{\max}
239 of epithelial phytase kinetics were found between Hubbard × Peterson and Ross × Ross
240 broilers, but not in pc InsP₆ hydrolysis (Applegate et al., 2003). It has been hypothesized from
241 genetic studies that epithelial phytase expression is affected by the bird's genetic background
242 (Beck et al., 2014). Indeed, genomic studies showed significant heritability in the range of
243 0.10 to 0.22 for P utilization, P excretion rate, and phytate-P bioavailability in broilers and
244 Japanese quail (Beck et al., 2016; de Verdal et al., 2011; Zhang et al., 2003). In the current
245 ring test, fast growing broilers from different strains were used that were typical for the
246 respective station. It is possible that the genetic background of the parent lines used for broiler
247 production in different regions of the world was different and this contributed to the
248 differences we found among stations. However, it is not currently possible to quantify the
249 contribution of epithelial phytase to intestinal luminal InsP₆ hydrolysis.

250 Some of the stations involved in the current study used unsexed broilers while others
251 used males. As a side aspect of the current study one station has compared males and females
252 when studying digestibility of the experimental diets (Schedle et al., 2016). The authors
253 reported almost identical values and no significant differences between males and females.

254 This let us conclude that for the current study it was not relevant whether mixed-sex or male
255 broilers were used.

256 Some other characteristics of the experiments such as growth, ADFI, age at sampling
257 and killing procedures were also different among stations. Statistical search for any
258 relationship would not be meaningful because the study was not designed to investigate any
259 of the effects and the data set is too limited for correlation analysis. However, it does not
260 appear from regressions that age at slaughter or at start of experimental feeding, BW at
261 slaughter, and ADFI in the trial period affected the results (Figure 5).

262
263 The protocol of WPSA (2013) has standardized several details for the determination of
264 P digestibility. From the results of this ring test, we conclude that the protocol standardization
265 must go beyond the standards already set. Differences in starter diet composition and
266 management conditions during the pre-experimental period of the trials affected the results
267 and thus also need standardization. Until a more standardized protocol is established, care
268 must be exercised when comparing P digestibility data from different laboratories.

269

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274

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Table 1. Composition of the experimental diets.

Diet	A	B	C
<i>Ingredients (g/kg)</i>			
Soybean meal from de-hulled seed ¹	400.0	510.0	620.0
Corn starch	448.6	336.6	224.6
Limestone (finely ground)	7.4	9.4	11.4
Soybean oil	30.0	30.0	30.0
Dried egg albumen	18.0	18.0	18.0
Sucrose	80.0	80.0	80.0
Sodium chloride	3.0	3.0	3.0
DL-Methionine	2.7	2.7	2.7
L-Threonine	0.3	0.3	0.3
Titanium dioxide	5.0	5.0	5.0
Vitamin and trace element premix ²	5.0	5.0	5.0
<i>Analyzed (g/kg, on dry matter basis)</i>			
CP	231	288	339
Total P ³	3.02 (0.06)	3.80 (0.08)	4.59 (0.08)
Ca ³	4.57 (0.09)	5.65 (0.09)	6.78 (0.13)
Ti ³	3.20 (0.07)	3.18 (0.07)	3.17 (0.07)
InsP ₆ -P	1.63	1.97	2.28
Ins(1,2,4,5,6)P ₅ -P ⁴	0.24	0.27	0.32

¹ The soybean meal contained per kg of dry matter (per analyses): 541 g crude protein, 76 g ash, 36 g crude fat, 41 g crude fiber, 116 g neutral detergent fiber, 72 g acid detergent fiber, 7.1 g P, 4.3 g InsP₆-P, 3.3 g Ca.

² Premix provided the following (per kilogram of diet): vitamin A, 12,000 IU; cholecalciferol, 63 µg; vitamin E, 50 IU; vitamin K₃, 1.5 mg; vitamin B₁, 2.0 mg; vitamin B₂, 7.5 mg; vitamin B₆, 3.5 mg; vitamin B₁₂, 20 mcg; niacin, 35 mg; D-pantothenic acid, 12 mg; choline chloride, 460 mg; folic acid, 1.0 mg; biotin, 0.2 mg; Fe, 80 mg (as FeSO₄·H₂O); Cu, 12 mg (as CuSO₄·5H₂O); Mn, 85 mg (as MnO); Zn, 60 mg (as ZnSO₄·H₂O); Co, 0.4 mg (as CoSO₄·7H₂O); I, 0.8 mg (as KI); Se, 0.15 mg (as Na₂SeO₃).

³ Mean (standard deviation) of analyses made for each laboratory.

⁴ All other lower inositol phosphate isomers were below the limit of quantification.

Table 2. Summary of the broiler chicken trials conducted in the collaborating stations.

Station number	Broiler strain	Sex	No of experimental replicates per diet	Birds per replicate	Age when placed on treatment diets (d)	Age at sampling (d)	BW before sampling (kg)	ADFI (g)	Feed/gain during the experimental period	Method of killing
1	Ross 708	male	8	8	20	25	1.12	123	1.63	CO ₂ asphyxiation
2	Ross PM3	unsexed	8	8	23	28	1.86	174	1.22	Pentobarbital injection
3	Ross 308	unsexed	6 ¹	8	19	27	1.46	138	1.32	Cervical dislocation
4	Ross 308	unsexed	6	8	22	28	1.84	163	1.27	Cervical dislocation
5	Ross 308	male	6	10	22	27	1.03	81	1.26	CO ₂ asphyxiation ²
6	Ross 308	unsexed	6	8	23	28	1.52	151	1.06	Cervical dislocation
7	Ross 308	male	8	12	17	22	1.16	119	1.36	Cervical dislocation
8	Ross 308	male	6	12	20	27	1.57	133	1.35	Pentobarbital injection
9	Heritage 5632	male	6	8	16	21	0.82	93	1.43	CO ₂ asphyxiation
10	Ross 708	unsexed	6	8	20	25	1.11	109	1.44	CO ₂ asphyxiation
11	Ross 308	unsexed	7	8	18	23	1.03	93	1.42	CO ₂ asphyxiation
12	Ross 308	male	8	8	16	25	1.45	98	1.32	Stunning
13	Ross PM3	unsexed	10	8	14	21	1.02	103	1.42	Cervical dislocation
14	Ross PM3	male	6 ¹	8	17	24	1.3	128	1.37	Pentobarbital injection
15	Ross 308	unsexed	6	10	18	28	1.72	140	1.40	Cervical dislocation
16	Ross PM3	unsexed	10 ¹	4	12	22	1.00	70	1.28	CO ₂ asphyxiation
17	Ross 308	male	6	12	14	24	1.24	96	1.31	Sedanum and Ketamin

¹Each replicate comprised two pooled cages.²CO₂ asphyxiation following anesthesia with a gas mixture of 35% CO₂, 35% N₂, and 30% O₂.

Table 3. Prececal digestibility of P and Ca of diets with variable inclusion of soybean meal and limestone as determined in 16 trials with broiler chickens

Diet	No of station	n ²	P digestibility ¹ (%)						Ca digestibility ¹ (%)					
			A		B		C		A		B		C	
			Mean	SD	Mean	sd	Mean	SD	Mean	SD	Mean	SD	Mean	SD
1		8	58	2.7	52	5.0	48	4.9	47	8.9	47	6.6	39	8.6
2		8 ³	67	3.2	58	3.3	52	1.2	64	3.4	55	5.6	47	3.3
3		6	68	4.8	62	2.2	50	3.2	56	5.0	49	3.1	42	3.4
4		6 ⁴	63	3.9	56	3.7	52	2.1	65	9.2	49	4.8	44	3.5
5		6	73	1.9	69	2.9	66	2.9	62	2.5	58	2.3	54	3.8
6		6	61	3.8	60	4.1	53	7.2	55	7.5	51	11.6	41	7.4
7		8	59	5.3	54	3.2	50	4.0	50	4.9	44	4.8	39	3.4
8		6	71	2.2	62	4.9	62	3.6	62	2.9	53	5.5	54	4.2
9		6	62	4.4	55	2.9	48	3.9	55	7.5	45	5.6	39	6.6
10		6	68	2.5	61	2.9	53	1.5	58	4.0	52	6.5	43	3.7
11		7	55	5.5	46	5.3	45	6.2	56	6.3	46	5.1	43	5.7
12		8	71	4.0	61	4.2	55	5.2	64	4.3	55	3.8	49	4.6
13		10	72	5.6	70	6.3	62	2.7	58	5.8	55	9.7	51	6.4
15		6	74	2.3	63	9.7	61	6.9	62	1.7	51	8.7	50	7.8
16		10	72	2.8	66	4.9	59	4.9	44	5.7	45	7.6	43	7.7
17		6	82	1.6	79	3.2	69	2.4	71	3.4	68	2.7	58	4.0

Diets: A = 400 g/kg soybean meal; B = 525 g/kg soybean meal; C = 650 g/kg soybean meal.

¹Effects of diet, station, and their interaction were statistically significant ($P < 0.05$).

²Number of replicates (cages) per diet. ³n = 7 for diet C. ⁴n = 5 for diets A and C.

Table 4. Results of linear regression analysis of prececal digestible P concentration (y, g/kg DM) in function of dietary P concentration (x, g/kg DM) using diets with incremental inclusion levels of soybean meal

No of station	Intercept	<i>SE</i>	Slope		<i>SE</i>	R ²	Root MS error
1	0.83	0.30	0.31	^{abc}	0.07	>0.99	0.01
2	1.37	0.41	0.22	^{bc}	0.08	0.97	0.02
3	1.50	0.43	0.19	^c	0.07	0.68	0.09
4	0.99	0.40	0.30	^{abc}	0.07	0.98	0.03
5	0.66	0.42	0.51	^a	0.08	>0.99	<0.01
6	0.72	0.42	0.38	^{abc}	0.08	0.93	0.07
7	0.79	0.42	0.33	^{abc}	0.08	>0.99	<0.01
8	0.80	0.42	0.43	^{ab}	0.07	0.95	0.06
9	1.34	0.42	0.19	^c	0.08	0.91	0.04
10	1.36	0.43	0.25	^{bc}	0.07	0.94	0.04
11	0.81	0.42	0.27	^{bc}	0.08	0.92	0.05
12	1.34	0.42	0.26	^{bc}	0.07	>0.99	0.02
13	0.94	0.42	0.42	^{ab}	0.07	0.95	0.07
15	1.02	0.41	0.38	^{abc}	0.07	0.97	0.04
16	1.12	0.41	0.34	^{abc}	0.08	>0.99	0.02
17	1.37	0.42	0.41	^{abc}	0.08	0.91	0.08

Superscript letters indicate significant differences between slopes.

Table 5. Prececal disappearance of InsP₆ of diets with variable inclusion of soybean meal as determined in 4 trials with broiler chicken

Diet	No of station	Prececal InsP ₆ disappearance ¹ (%)					
		A		B		C	
	n ²	Mean	SD	Mean	SD	Mean	SD
4	6 ³	70	4.5	52	8.5	47	5.0
5	6	74	3.8	67	4.7	58	7.1
7	8	58	10.8	47	8.1	36	8.9
9	6	62	4.3	43	3.6	23	11.4

Diets: A = 400 g/kg soybean meal; B = 525 g/kg soybean meal; C = 650 g/kg soybean meal.

¹Effects of diet, station, and their interaction were statistically significant ($P < 0.05$).

²Number of replicate cages.

³n = 5 for diets A and C.

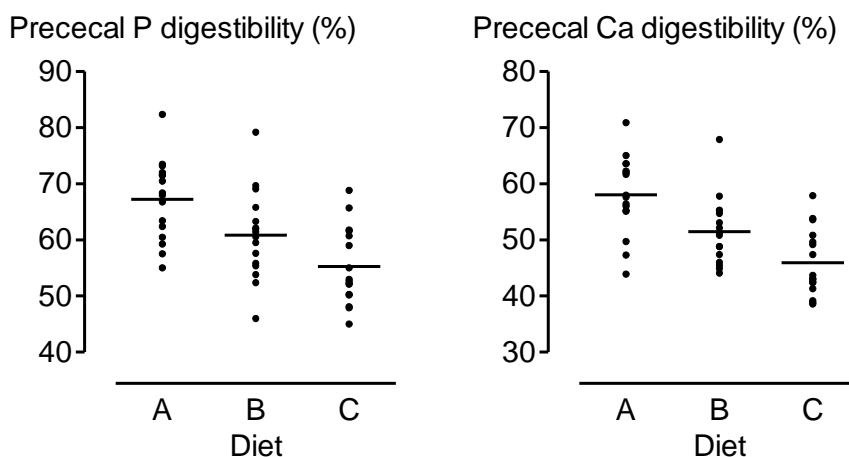


Figure 1. Scatter plot of prececal digestibility data from 16 broiler chicken trials conducted in 16 stations and using diets containing 400 (A), 525 (B) or 650 (C) g/kg soybean meal (each dot shows the mean of one station for the respective diet).

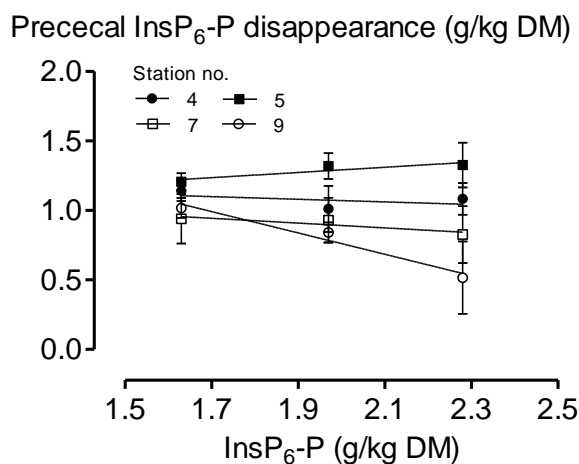


Figure 2. Relationship between the concentration of InsP₆-P in the diet and prececal InsP₆-P disappearance in diets with different levels of soybean meal and studied in 4 broiler trials in 4 stations (means and SD). Slopes estimated for stations 4, 5, and 7 did not deviate from zero ($P > 0.05$).

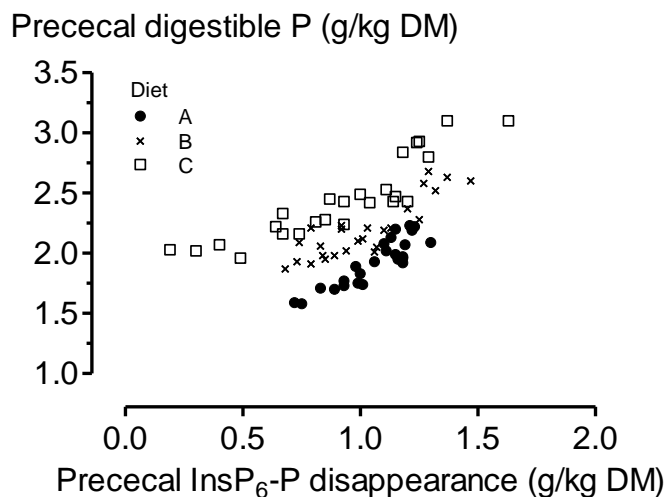


Figure 3. Relationship between the concentration of prececal digestible P in dependence of prececal InsP₆-P disappearance in diets including different levels of soybean meal and studied in 4 broiler trials in 4 stations (each symbol shows the value of one cage). The slope of the linear regressions calculated for each diet were not significantly different ($P = 0.27$), and the pooled slope was 0.91.

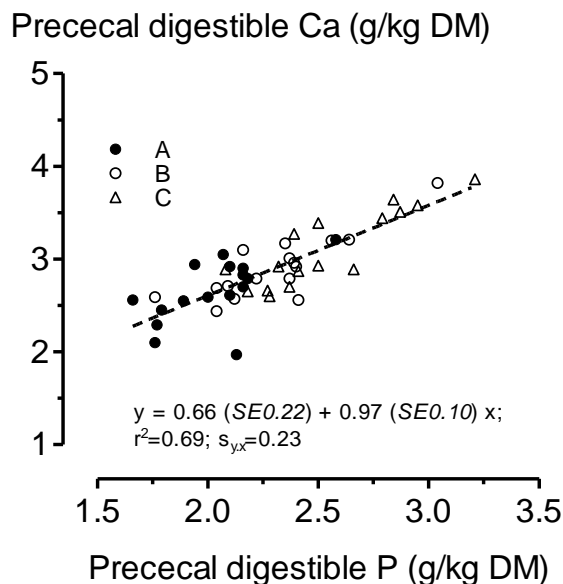


Figure 4. Relationship between the concentration of prececal digestible P and prececal digestible Ca in diets fed to broilers and using diets containing 400 (A), 525 (B) or 650 (C) g/kg soybean meal (each symbol shows the mean of one out of 16 stations for the respective diet). The slopes determined separately for each diet were not significantly different.

Figure 5. Comparison between the prececal digestibility of P in diet A with the formulated P concentration of the pre-experimental starter feed (panel A), the supplemented phytase of the pre-experimental starter feed (panel B), the age of birds when the experimental diets were introduced (panel C), the duration of the experimental feeding period (panel D), the age of birds at the day of sampling (panel E), the BW of the birds at the day of sampling (panel F), and the ADFI in the trial period (panel G). Each dot shows the mean of one station. The difference to zero of the calculated slopes of linear regressions was significant in panel D ($P < 0.01$), but not in panel A ($P = 0.08$) or any other panel ($P > 0.1$).