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(Article begins on next page)



Effects of dietary *Lactobacillus acidophilus* and *Bacillus subtilis* on laying performance, egg quality, blood biochemistry and immune response of organic reared laying hens.

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1 Effects of dietary *Lactobacillus acidophilus* and *Bacillus subtilis* on laying performance, egg
2 quality, blood biochemistry and immune response of organic reared laying hens.

3

4 Short title: Effects of probiotics on organic reared hens

5

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13

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15

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Comment [FB2]: Key words (give 4-5 key words and arrange a-z), separate by, and each entry starts with first capital Word

40 INTRODUCTION

41

42 In recent years, attention to nutrition-based health strategies has grown (Trovato, 2012), and
43 because increased microbial resistance to antibiotics and residues in animal products can be harmful
44 to consumers, the interest in alternative strategies to reduce antibiotic use in animal production has
45 increased.

46 A probiotic is a non-pathogenic and non-toxic microorganism (e.g., bacteria, fungi and others)
47 administered through the digestive tract. The FAO/WHO (2001) defines probiotics as “live
48 microorganisms which, when administered in adequate amounts, confer a health benefit on the
49 host”. The primary microorganisms used as probiotics in poultry nutrition are bacteria, such as
50 *Lactobacillus* spp., *Enterococcus* spp., *Pediococcus* spp., *Streptococcus* spp., *Bifidobacterium* spp.
51 and *Bacillus* spp., as well as yeasts, such as *Saccharomyces cerevisiae* and *Candida* spp. (Ayasan,
2013).

52 The organic farming of laying hens has become an important economic activity in many countries
53 due partly to increasing environmental awareness and partly to increasing consumer demand for
54 “natural” products (Berg, 2001). In organic farming, the use of pharmacological products is strictly
55 controlled, so it becomes extremely important to ensure a high level of well-being for the animal, an
56 appropriate level of health and an effective defence against external *noxae* arising from ground
57 breeding. Probiotics could help to ensure such results without breaking the rules of organic farming.
58 The introduction of probiotics to the diet of laying hens can result in many positive outcomes:
59 increased feed consumption and improved feed conversion ratio (Nahashon et al., 1994; Haddadin
60 et al., 1996), higher resistance to parasitic infections (Dalloul et al., 2003; Tierney et al., 2004),
61 increased immune function (Dalloul et al., 2003), larger eggs (Davis and Anderson, 2002), higher
62 egg production and quality (Kurtoglu et al., 2004; Gallazzi et al., 2009; Panda et al., 2008),
63 decreased presence of cholesterol in the eggs (Mohan et al., 1995; Abdulrahim et al., 1996;
64 Kurtoglu et al., 2004; Zhang and Kim, 2013), decreased ammonia emissions (Zhang and Kim,

Comment [FB4]: The below mentioned literature is related to this paper, but authors did not use them in the text. It must be added these references.

AYASAN T, 2013. Effects of dietary inclusion of protexin (probiotic) on hatchability of Japanese quails. *Indian J Anim Sci*, 83(1): 78-81.

65 2013), reduced serum cholesterol (Fathi, 2013; Aluwong et al., 2012) and reduced oxidative stress
66 (Anwar et al., 2012).

67 In this study, two microorganisms, which differ in their ecology and metabolism, were selected:
68 *Lactobacillus acidophilus* D2/CSL, a commensal bacterium normally present in the intestinal
69 microbial population, and *Bacillus subtilis* (ATCC PTA-6737), a non-commensal, spore-forming
70 bacterium. Both bacterial species are included in the European Union Register of Feed Additives
71 (2013) pursuant to Regulation (EC) 1831/2003. The aim of this research was to investigate the
72 effects of probiotic dietary supplementation on production parameters, immune function, oxidative
73 stress and biochemical chemistry parameters of organically farmed laying hens.

Comment [FB5]: Is the problem significant and concisely stated?: Yes

74 MATERIALS AND METHODS

75 *Birds, feeding and management*

76 The study was conducted at a farm located in the Lazio region of Italy. A total of 900 16-week-old
77 Hy-Line layer hybrids were reared under organic farming guidelines according to the Council
78 Regulation (EC) No 1804/1999. The hens were randomly assigned to three groups of 300 birds
79 each, which were each divided into three pens (experimental units) of 100 birds each. The hens
80 where reared for a total of 20 weeks and received a pre-deposition diet for the first 4 weeks of the
81 experiment and the same deposition diet for the rest of the experimental period.

82 The control (CTR) group was fed a corn-soybean cake-based diet (Table 1). The L group was fed
83 the same diet supplemented with 0.1% *Lactobacillus acidophilus* (Lactomalt D2 Bio®,
84 *Lactobacillus acidophilus* D2/CSL CECT 4529, Zoo Assets Srl, Mantova, Italy) while the B group
85 was fed the same diet supplemented with 0.05% *Bacillus subtilis* (Clostat® brand dry – 740210,
86 *Bacillus subtilis* PB6 ATCC-PTA 6737, Kemin®, Herentals, Belgium). The experimental diets and
87 water were given to the birds *ad libitum*.

88 The birds were managed under the same prophylactic procedures. Throughout the experiment, the
89 natural photoperiod, temperature and humidity were maintained.

90

91 *Feed analysis*

92 The chemical composition of the diets was determined according to AOAC methods (2000, 1990,
93 1996). Starch content was evaluated according to ISO 10520:1997, and metabolisable energy
94 content was estimated using the equation of Carpenter and Clegg (1956).

95

96 *Performance*

97 Data for calculating the deposition rate, feed intake and feed conversion efficiency (FCE) were
98 recorded per pen (3 pens/group) daily at the beginning (weeks 5 and 6: T1) and end (weeks 19 and

Comment [FB6]: With which nutrient recommendation guide and with which diets formulation software program?

Comment [FB7]: Give information

Comment [FB8]: How was calculated????

99 20: T2) of the experiment after the hens reached the age of 30 and 58 weeks, respectively. Thirty
100 eggs/pen were collected for analysis at both T1 and T2.

101

102 *Chemical and physical analysis of eggs*

103 The chemical composition of the yolk was determined according to the AOAC (1995). Total protein
104 was calculated from Kjeldahl nitrogen using 6.25 as the conversion factor, and total lipids were
105 twice extracted from 5 g of each sample homogenate and separated by gravity (Folch et al., 1957).

106 Data recorded included the integrity, weight and thickness of the shell (Mueller and Scott, 1940), the
107 weight and colour of the yolk (Roche scale), and the height of the albumen (Haugh unit) using an
108 electronic gauge (Bukley et al., 1981). Colour coordinates (Commission International de l'Eclairage,
109 1976) were determined using a Minolta Chromameter CR400 (Minolta, Osaka, Japan — D65 light
110 source calibrated against a standard white tile), and the results were expressed in terms of lightness
111 (L*), redness (a*), and yellowness (b*).

112

113 *Serum analysis*

114 Fifteen hens/pen were sampled using vacuum tubes without an anticoagulant (Vacuette, Greiner
115 Bio-one, Frickenhausen, Germany) from the brachial vein on the first day of each sampling period
116 (T1 and T2). Blood samples were incubated at room temperature for 1 h and centrifuged at 1200 g
117 for 10 minutes. The serum samples were stored at -80°C to determine the metabolic profile, innate
118 immune parameters (lysozyme, serum bactericidal activity, and haemolytic complement assay),
119 oxidative stress and immune response to NDV antigens.

120 *Metabolic profile:* Serum samples were tested for ALT, AST, γ GT, cholesterol and triglycerides.

121 These tests were assessed by a Konelab 2001 biochemical analyser using specific kits (Sentinel

122 Diagnostics, Milan, Italy).

Comment [FB9]: Are the methods described comprehensively?: Yes

Comment [FB10]: Ethical approval for the research was not informed in the manuscript.

The authors must include the Protocol number of the Ethics Committee of Animal Use or explain why this information wasn't include in this paper, since the authors used layer for this experiment.

Comment [FB11]: Give relevant references

Comment [FB12]: Give relevant references

123 *Serum bactericidal activity (SBA)*. The SBA was performed according to a method previously
124 validated for cattle (Amadori et al., 1997). The test is based on a serum challenge with non-
125 pathogenic *E. coli*, and its concentration is expressed as a percentage.

126 *Lysozyme*. Serum lysozyme was measured with a lysoplate assay (Osserman and Lawlor, 1966)
127 carried out in a moist incubator at 37°C for 18 minutes. The method is based on the *Micrococcus*
128 *lysodeikticus* lyses in 1% agarose. The diameter of the lysed zones was measured with a ruler and
129 compared with the lysed zones of a standard lysozyme preparation (Sigma, Milan, Italy, M 3770),
130 and the value is expressed in µg/mL.

131 *Haemolytic complement assay (HCA)*. The haemolytic complement assay (Barta and Barta, 1993)
132 was carried out on microtitre plates. The complement titre is the reciprocal of the serum dilution
133 that causes 50% of ram red blood cells to lyse, and its concentration is expressed as CH₅₀%.

134 *Haemagglutination inhibition test for NDV (HI)*. Hens were vaccinated at the hatchery against
135 Newcastle disease. The test to determine the production of NDV antibodies is based on the
136 principle that the haemagglutinin on the viral envelope can bring about the agglutination of chicken
137 red blood cells and that this can be inhibited by specific antibodies. V-bottomed microtitration
138 plates were used, and this test was performed according to the OIE Manual (2012). The HI titre was
139 expressed as the log₂, reciprocal of the highest serum dilution producing 10% inhibition of HI
140 activity.

141 *Oxidative status*. The reactive oxygen substances (ROS) in the serum were evaluated with a
142 commercial kit (Diacron, Grosseto, Italy) and are expressed as mmol H₂O₂.

143 The serum antioxidant power (AP) was measured with a commercial kit (Diacron, Grosseto, Italy)
144 that evaluates the ability of plasma to oppose the oxidative action of a hypochlorous acid (HClO)
145 solution. The AP levels of the sample are expressed as µmol of neutralised HClO.

146

147 *Statistical analysis*

Comment [FB13]: Give relevant references

148 The data were analysed using the GLM procedure in SAS (2010). The ANOVA model included the
149 diet (C, L, and B) and sampling time (T1 and T2) as fixed factors, as well as their interaction. Data
150 are reported as least squares means \pm standard error. Differences were assessed by Tukey's test and
151 considered to be significant when $P < 0.05$.

152

153 RESULTS

154 *Hen productive performance*

155 Productive performance of laying hens is shown in Table 2. No differences between dietary groups
156 or sampling time were observed ($P > 0.05$). At T2, an increase in egg size was recorded with the
157 increased age of the hens.

158

159 *Hen biochemical profiles*

160 The results for the clinical chemistry parameters are reported in Table 3. All parameters except
161 GGT were affected by the diet, and the cholesterol values of both of the supplemented groups were
162 lower ($P < 0.001$) at T2. The triglyceride content of serum was only lower ($P < 0.05$) at T2 for the B
163 group.

164 As for the enzymes, ALT followed a similar trend; the values of the treated groups were lower
165 ($P < 0.001$) than the control group at T2. The lowest AST levels were recorded in the B group at T1.

166

167 *Immune response*

168 The results of the innate immunity profiles are reported in Table 4. The bactericidal activity of the
169 blood was increased at T2 by both of the probiotic supplementations. Lower HC values ($P = 0.008$)
170 were observed for the L group compared to the C group (overall effect), and the lysozyme
171 concentration was lower ($P = 0.016$) for the B group at T1. An overall effect of the L diet on the
172 production of antibodies ($P = 0.013$) against NDV was observed, but these values were not affected
173 by the age of the birds.

Comment [FB14]: does the RESULTS section provide adequate presentation of the authors' own results?
YES

174

175 *Oxidative status*

176 Oxidative status data are reported in Table 5. No differences between the dietary groups were
177 recorded.

178

179 *Egg physicochemical characteristics*

180 No effects of diet on egg quality were observed (Table 6). Among the physical characteristics of
the eggs,

181 only the Roche scale colour was affected ($P<0.05$) by diet; supplementation with 0.1%

182 *L. acidophilus* produced paler egg yolks compared to both the C and B groups.

183 Sampling time had the greatest effect on the egg physical characteristics. Yolk weight, albumen

184 weight, shell weight, yolk percentage, albumen percentage, edible percentage, Roche scale, Haugh

185 unit and colour (a and b values) all increased ($P<0.05$) at T2.

186 The egg chemical characteristics (Table 7) were almost unaffected by supplementation with

187 probiotics, except for the lipid content which was decreased ($P<0.05$) in the L group.

188

189 DISCUSSION

190 *Hen productive performance*

191 The overall results are similar to those found in other studies aimed at investigating the productivity

192 of laying hens under organic conditions (Hammershoj and Steenfeldt, 2005; Mugnai et al., 2009;
Minelli et al., 2007; Hammershoj and

193 Steenfeldt, 2005). According to Cunningham et al. (1960) and Funk and Kempster (1934),

194 physiological ageing, with the consequent reduction in productive performance, increases egg size,

195 and previous studies have shown that dietary probiotics, such as *Lactobacillus* spp. and *Bacillus*

196 spp., could increase the performance of laying hens (Zhang and Kim, 2013; Xu et al., 2006;

197 Abdulrahim et al., 1996; Nahashon et al., 1994). In contrast, other studies did not record any

198 positive effects (Mikulski et al., 2012; Tortuero and Fernandez, 1995; Nahashon et al., 1996), but

199 the values obtained in this study are difficult to compare with those data because they were not

Comment [FB15]: Discussion section is not enough to properly explain reasons of the results obtained

Comment [FB16]: Avoid old references.

Comment [FB17]: Avoid old references.

Comment [FB18]: On the year basis

200 obtained under organic conditions. Other reasons for the discrepancies between studies could be the
201 age of the study animals as well as the species and the dose of microorganisms used.

202

203 *Hen biochemical profiles*

204 Regarding lipid metabolism, the reduction in circulating cholesterol observed in the probiotic-
205 treated groups in this study agrees with the results of other researchers (Onifade et al., 1999;
206 Jouybari et al., 2009). Fathi (2013) found that supplementation with *Lactobacillus cultures*
207 decreased serum cholesterol and triglyceride levels, and similar results were reported by Panda et al.
208 (2006) who found that serum total cholesterol and triglycerides were reduced by dietary
209 supplementation with *Lactobacillus sporogenes*. Hashemzadeh et al. (2013) found reduced serum
210 cholesterol in groups treated with *Lactobacillus rhamnosus*. Probiotics could contribute to the
211 regulation of serum cholesterol concentrations through the deconjugation of bile acids, and because
212 the excretion of deconjugated bile acids is enhanced, more cholesterol molecules could be spent for
213 the recovery of bile acids (De Smet et al., 1994). Other authors have ascribed this result to the
214 reduced adsorption and/or synthesis of cholesterol in the gastrointestinal tract (Mohan et al., 1995; [add another
reference](#)),

215 but it has also been speculated that *L. acidophilus* could be able to decrease the cholesterol in the
216 blood through the deconjugation of bile salts in the intestinal lumen thereby preventing them from
217 acting as precursors in cholesterol synthesis (Abdularahim et al., 1996).

218 The trend observed for ALT and AST levels confirms the data presented by Fathi (2013), who
219 reported that *Lactobacillus* supplementation can decrease ALT and AST plasma concentrations. On
220 the contrary, other authors ([Baidya et al., 1994](#); Capcarova et al., 2011; ~~[Baidya et al., 1994](#)~~) found that ALT and
AST

221 were not affected by the dietary addition of probiotic strains. In rats, however, dietary *Lactobacillus*
222 *plantarum* and *Bifidobacterium infantis* decreased the level of ALT (Osman et al., 2007).

223

224 *Immune response*

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225 Probiotics are non-pathogenic bacteria that can promote animal health by reducing pathogen
226 colonisation (Mead, 2005); these reductions are attributed to competitive exclusion, increased
227 volatile fatty acid production, and the potentiation of the immune system (Nava et al., 2005,
228 Donoghue et al., 2006). Fong et al. (2015) observed an improved immune response in the cells of
229 healthy subjects treated with *Lactobacillus ramosus* in vitro, and lactic acid bacteria can markedly
230 increase the humoral immune response in chickens (Koenen et al., 2004).

231 The bactericidal property of blood may be due to the presence of complementary factors and a
232 small quantity of natural antibodies (Michael et al., 1962) as well as other antibacterial substances,
233 such as beta-lysine and lectin (Donaldson et al., 1964; Kawasaki et al., 1989). Our results showed
234 an increase in serum bactericidal activity in T2 for both probiotic supplements. It is possible to
235 assume that probiotics, which contribute to the maintenance of the integrity of the intestinal
236 membrane, are able to enhance intestinal immunity (i.e., higher levels of IgA; Gorbach, 2000) and
237 could be able to facilitate the intestinal absorption of antibacterial substances that are able to
238 improve the AP parameters.

239 Unexpectedly, lower complementary values were registered for the L group compared to the C
240 group in our study, and our data do not allow for a reasonable explanation of this trend.

241 Lysozyme is able to damage bacterial cell walls by attacking peptidoglycan. Because this enzyme is
242 largely present in the granules of several types of cells, its serum concentration can provide useful
243 information about granulocyte activities and the functionality of the monocyte-macrophage system;
244 therefore, it can be reasonably considered to be a possible marker for the quantification of
245 pathogens in the environment (Gordon et al., 1974). In this study, the concentration of lysozyme
246 was influenced by the use of probiotics at T1, and it was lower in the B group compared to control,
247 which is a sign of lower degranulation of granulocytes. In their study on rats, Vilahur et al. (2014)
248 showed that by modulating the innate immune response, dietary *Lactobacillus plantarum* could be a
249 natural option to protect against inflammatory disorders.

Comment [FB19]: WHY OLD REFERENCE,
why???

250 As for the immune response to the NDV vaccine, the L group showed an antibody concentration
251 higher than that observed in the control group in accordance with other studies (Khaksefidi and
252 Ghoorchi, 2006). This difference in antibody production could be attributed to the probiotic
253 microorganism, which might have influenced the intestinal ecosystem and aided the assimilation of
254 nutrients essential for triggering the immune cells to produce antibodies (Panda et al., 2000).

255

256 *Oxidative status*

257 Probiotics have shown antioxidant properties in poultry reared under intensive conditions
258 (Capcarova et al., 2010), but in the present study, no differences between groups were found in the
259 oxidative parameters evaluated. We can hypothesise that the antioxidant properties of probiotics
260 found by other authors might become evident in stressful environments, and the more “natural”
261 organic farming conditions may have prevented this dietary supplementation from producing any
262 favourable effects for animal metabolism.

263

264 *Egg physiochemical characteristics*

265 With respect to egg physical parameters (Table 3), probiotics have been reported to increase egg
266 quality (decreased yolk cholesterol level, improved shell thickness and egg weight -Kurtoglu et al.,
267 2004; Xu et al., 2006). The results obtained in the present study are in accordance with Yörük et al.
268 (2004) who found no effect of diet on egg quality.

269 In terms of albumen quality, Williams (1992) found that, excluding disease, Haugh unit scores
270 decrease with advancing flock age, but they increased in our study.

271 The values registered for egg lipid content (Table 7) are in accord with the observed reduction in
272 serum total cholesterol and triglycerides. Egg cholesterol content showed a time-dependent effect
273 ($P<0.05$), which is considered to be physiological; with an increasing deposition period, hens
274 reduce their productive performance while cholesterol concentration increases (dilution effect).

275 These results are in agreement with Minelli et al. (2007), who found lower percentages of lipids in
276 organic eggs collected at the end of the deposition cycle.

277

278 CONCLUSIONS

279 In this study, dietary supplementation with two probiotic microorganisms, *L. acidophilus* D2/CSL
280 CECT 4529 and *B. subtilis* PB6 ATCC-PTA 6737, resulted in changes to the blood chemical
281 parameters and immune response of organically reared hens. In particular, *L. acidophilus* and *B.*
282 *subtilis* induced positive changes in metabolic parameters, such as serum cholesterol and
283 triglyceride content and ALT values, without changing the productive performance, egg
284 physiochemical characteristics and oxidative status of the birds. The administration of *L.*
285 *acidophilus* resulted in an improvement in antibody production against the Newcastle disease virus.
286 In conclusion, these probiotic strains could be used in organic farming to improve animal welfare
287 and immune defence.

288

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Comment [FB20]: Are the interpretations and conclusions justified by the results?: Yes

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Comment [FB21]: Uniformity in references must be seen.

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451 density diets on performance, egg quality, excreta microflora, excreta noxious gas emission, and
452 serum cholesterol concentrations in laying hens. *Journal of Animal Science* **91**, 4781-4787.

For Peer Review

1 Table 1. Ingredients and chemical composition of the experimental diets
2
3
4

	Predeposition	Deposition
<i>Ingredients (%)</i>		
Maize	55.00	43.50
Soybean cake	25.00	30.00
Wheat bran	7.00	7.00
Broadbean	5.00	
Wheat		5.00
Calcium carbonate	3.00	8.70
Soybean	2.50	3.00
Dicalcium phosphate	1.50	1.50
Sodium chloride	0.30	0.50
Sodium bicarbonate	0.20	0.30
Mineral and Vitamin Premix*	0.50	0.50
<i>Calculated composition</i>		
M.E. (kcal/kg) †	3.041	2.810
<i>Analysed composition (%)</i>		
Moisture	8.28	8.35
Protein	16.92	18.01
Lipids	4.61	5.48
Ash	8.51	11.48
NDF	13.85	13.51
ADF	3.92	3.84
ADL	0.79	0.72
Ca	2.94	3.88
P	0.62	0.62
Lys	0.89	1.01
Met	0.45	0.36
Na	0.17	0.16

5 *Vitamin and mineral premix supplied per kilogram of diet: Vitamin A, 14000 UI; Vitamin D3,
6 3000 UI; Vitamin E, 30.00 mg, Vitamin K, 2.00 mg; Vitamin B₁, 1.75 mg; Vitamin B₂, 12.00 mg;
7 Vitamin B₆, 2.00 mg; Vitamin B₁₂, 0.015 mg; Niacin, 35.00 mg; Folic acid, 0.50mg; Pantothenic
8 acid, 10.00 mg; Fe (FeCO₃), 44.51 mg; Cu (CuSO₄.5H₂O), 58.95 mg; Mn (MnO), 193.5 mg; Zn
9 (ZnO), 69.75 mg; Se (Na₂SeO₃) 66.00 mg

10 †Metabolizable Energy content of diets was estimated using the equation of Carpenter and Clegg
11 (1956).

12 Table 2. Effect of probiotic dietary supplementation and sampling time on performance of laying
 13 hens

14

	T1	T2	Overall	Main effect	P
Deposition rate (%)					
C	69.82	79.15	76.05	Diet	0.062
B	70.90	80.50	77.09	Time	0.069
L	71.10	80.81	77.15	Diet x Time	0.064
SEM	2.897	3.215	2.954		
Feed efficiency					
C	2.54	2.71	2.63	Diet	0.075
B	2.50	2.68	2.58	Time	0.062
L	2.47	2.66	2.56	Diet x Time	0.069
SEM	0.181	0.215	0.205		
Egg weight (g)					
C	57.81	62.56	60.18	Diet	0.948
B	56.12	63.93	59.83	Time	<0.001
L	56.37	63.33	59.85	Diet x Time	0.500
SEM	1.063	1.344	1.157		

15
 16 N= 15 hens pen/group per time (T1 and T2).

17 ^{A,B...} Means within a column lacking a common superscript differ (P<0.05)

18 C: basal diet; L: basal diet + 0.1 % of *Lactobacillus acidophilus*; B: basal diet + 0.05% of *Bacillus*
 19 *subtilis*.

20 Table 3. Effect of probiotic dietary supplementation and sampling time on clinical chemistry
 21 parameters of laying hens
 22
 23

	T1	T2	Overall	Main effect	P
Cholesterol (mmol/l)					
C	2.86	2.85 A	2.86 A	Diet	<0.001
B	2.41	2.27 B	2.34 B	Time	0.329
L	2.42	2.21 B	2.32 B	Diet x Time	0.793
SEM	0.151	0.143	0.106		
Triglycerides (mmol/l)					
C	9.39	10.97 A	10.18 A	Diet	0.002
B	7.29	7.75 B	7.88 B	Time	0.027
L	9.05	8.75 AB	9.12 AB	Diet x Time	0.368
SEM	0.543	0.583	0.397		
ALT (U/I)					
C	13.85 A	18.24 A	16.05 A	Diet	<0.001
B	12.03 A	9.97 B	11.00 B	Time	0.277
L	7.40 B	9.26 B	8.33 B	Diet x Time	0.013
SEM	1.230	1.954	1.069		
AST (U/I)					
C	205.39 A	169.05	187.22	Diet	0.064
B	162.09 B	174.65	168.37	Time	0.023
L	188.33 A	167.82	178.07	Diet x Time	0.009
SEM	7.222	8.121	5.427		
GGT (U/I)					
C	50.34	46.27	48.31	Diet	0.57
B	49.95	48.74	48.85	Time	0.7
L	48.69	50.73	49.71	Diet x Time	0.495
SEM	2.472	2.721	1.844		

24

25 N= 15 hens pen/group per time (T1 and T2).

26 ^{A,B}... Means within a column lacking a common superscript differ (P<0.05)27 C: basal diet; L: basal diet + 0.1 % of *Lactobacillus acidophilus*; B: basal diet + 0.05% of *Bacillus subtilis*.

28

Comment [FB26]: Ethical approval for the research was not informed in the manuscript.

The authors must include the Protocol number of the Ethics Committee of Animal Use or explain why this information wasn't include in this paper, since the authors used layer for this experiment.

29 Table 4. Effect of probiotic dietary supplementation and sampling time on immune response of
 30 laying hens
 31

	T1	T2	Overall	Main effect	P
SBA (%)					
C	42.77	43.97 B	43.37 B	Diet	0.010
B	43.36	58.16 A	50.76 AB	Time	<0.001
L	41.38	63.81 A	52.59 A	Diet x Time	0.004
SEM	3.161	2.963	2.257		
HCA (CH_{50%})					
C	27.56	31.53	29.54 A	Diet	0.008
B	22.46	31.89	27.17 AB	Time	0.003
L	21.83	24.16	22.99 B	Diet x Time	0.250
SEM	1.863	2.621	1.491		
Lysozyme (µg/mL)					
C	7.517 A	4.102	5.810	Diet	0.418
B	5.430 B	5.129	5.280	Time	<0.001
L	6.269 AB	4.067	5.168	Diet x Time	0.016
SEM	0.462	0.588	0.366		
NDV antibodies (HI titer in log₂)					
C	160.00	156.44	158.22 B	Diet	0.013
B	200.00	252.23	226.12 AB	Time	0.455
L	277.33	310.86	294.09 A	Diet x Time	0.834
SEM	41.605	50.956	30.896		

32 N= 15 hens pen/group per time (T1 and T2).

33 ^{A,B...} Means within a column lacking a common superscript differ (P<0.05)

34 C: basal diet; L: basal diet + 0.1 % of *Lactobacillus acidophilus*; B: basal diet + 0.05% of *Bacillus subtilis*.

35
 36 SBA: serum bactericidal activity; HCA: haemolytic complement assay; NDV: Newcastle disease
 37 virus.

Comment [FB27]: Ethical approval for the research was not informed in the manuscript.

The authors must include the Protocol number of the Ethics Committee of Animal Use or explain why this information wasn't include in this paper, since the authors used layer for this experiment.

38 Table 5. Effect of probiotic dietary supplementation and sampling time on ROM's and AP
39

	T1	T2	Overall	Main effect	P
ROM's (mmol H ₂ O ₂)					
C	3.272	4.754	4.012	Diet	0.113
B	3.073	3.625	3.349	Time	0.078
L	3.115	3.087	3.101	Diet x Time	0.238
SEM	0.395	0.522	0.318		
AP (μmol/HClO)					
C	62.644	86.797	74.723	Diet	0.548
B	70.798	82.015	76.414	Time	0.891
L	85.765	77.111	76.442	Diet x Time	0.053
SEM	9.222	9.642	6.729		

40 N= 15 hens pen/group per time (T1 and T2).
 41 ^{A,B...} Means within a column lacking a common superscript differ (P<0.05)
 42 C: basal diet; L: basal diet + 0.1 % of *Lactobacillus acidophilus*; B: basal diet + 0.05% of *Bacillus*
 43 *subtilis*.
 44 | ROM's: reactive oxygen metabolites; AP: antioxidant power of serum

45 Table 6. Effect of probiotic dietary supplementation and sampling time on egg physical parameters
46

	T1	T2	Overall	Main effect	P
Yolk weight (g)					
C	13.58	16.38	14.98	Diet	0.192
B	13.42	16.14	14.78	Time	<0.001
L	13.33	15.70	14.51	Diet x Time	0.075
SEM	0.382	0.333	0.275		
Albumen weight(g)					
C	34.01	38.55	36.28	Diet	0.631
B	32.22	38.46	35.34	Time	<0.001
L	32.99	38.39	35.69	Diet x Time	0.697
SEM	0.794	1.310	0.701		
Shell weight(g)					
C	6.90	7.76	7.33	Diet	0.667
B	7.05	7.58	7.31	Time	<0.001
L	6.66	7.67	7.17	Diet x Time	0.469
SEM	0.199	0.163	0.141		
Shell ash (%)					
C	3.21	3.19	3.20	Diet	0.664
B	3.30	3.34	3.33	Time	0.082
L	2.94	3.53	3.23	Diet x Time	0.094
SEM	0.139	0.144	0.105		
Yolk (%)					
C	23.49	26.47	24.98	Diet	0.192
B	23.97	25.42	24.70	Time	<0.001
L	23.63	24.79	24.21	Diet x Time	0.075
SEM	0.412	0.379	0.301		
Albumen (%)					
C	58.77	62.13	60.45	Diet	0.411
B	57.46	60.47	58.96	Time	0.003
L	58.64	60.60	59.62	Diet x Time	0.810
SEM	0.631	1.730	0.789		
Shell (%)					
C	11.98	12.455	12.22	Diet	0.520
B	12.52	11.967	12.45	Time	0.723
L	11.81	12.129	11.97	Diet x Time	0.119
SEM	0.251	0.260	0.190		
Edible(%)					
C	82.26	88.60	85.43	Diet	0.204
B	81.43	85.90	83.66	Time	<0.001
L	82.27	85.39	83.83	Diet x Time	0.335

SEM	0.605	1.699	0.770		
Albumen/yolk					
C	2.53	2.38	2.45	Diet	0.582
B	2.43	2.39	2.41	Time	0.187
L	2.51	2.46	2.49	Diet x Time	0.690
SEM	0.059	0.092	0.0522		
Colour (Roche scale)					
C	5.72	8.32	7.02 A	Diet	0.007
B	5.82	7.92	6.85 A	Time	<0.001
L	4.83	7.30	6.06 B	Diet x Time	0.500
SEM	0.305	0.296	0.2260		
Hugh Unit					
C	120.08	123.95	122.02	Diet	0.125
B	120.74	124.41	122.58	Time	<0.001
L	118.25	123.90	121.08	Diet x Time	0.340
SEM	0.673	0.767	0.523		
Colour CIELAB					
L*					
C	56.11 B	57.81	56.96	Diet	0.095
B	57.24 AB	56.99	57.11	Time	0.175
L	57.86 A	58.01	57.93	Diet x Time	0.102
SEM	0.430	0.505	0.338		
a					
C	-4.41	-3.58	-3.96	Diet	0.078
B	-4.67	-3.52	-4.12	Time	<0.001
L	-5.01	-3.90	-4.46	Diet x Time	0.846
SEM	0.213	0.196	0.156		
b					
C	36.37	40.46	37.42	Diet	0.740
B	36.46	39.11	37.79	Time	<0.001
L	36.56	39.11	37.33	Diet x Time	0.018
SEM	0.612	0.513	0.438		

47 N=30 eggs/pen/group per time (T1 and T2).

48 A,B... Means within a column lacking a common superscript differ (P<0.05)

49 C: basal diet; L: basal diet + 0.1 % of *Lactobacillus acidophilus*; B: basal diet + 0.05% of *Bacillus subtilis*.

50

51 Table 7. Effect of probiotic dietary supplementation and sampling time in egg chemical parameters
 52

	T1	T2	Overall	Main effect	P
Ash (%)					
C	1.76	1.78	1.77	Diet	0.8975
B	1.78	1.78	1.78	Time	0.0407
L	1.73	1.81	1.77	Diet x Time	0.1168
SEM	0.019	0.018	0.014		
Crude protein (%)					
C	16.00	16.20	16.10	Diet	0.1404
B	15.87	16.02	15.95	Time	0.1001
L	16.12	16.29	16.20	Diet x Time	0.9835
SEM	0.109	0.148	0.090		
Lipid (%)					
C	27.74 A	27.45	27.59 A	Diet	0.0009
B	27.70 A	27.56	27.63 A	Time	0.8460
L	26.80 B	27.31	27.05 B	Diet x Time	0.0407
SEM	0.138	0.193	0.115		
Cholesterol (mg/g yolk)					
C	12.80	12.52	12.66	Diet	0.6866
B	13.21	13.09	13.15	Time	0.7356
L	12.91	12.85	12.88	Diet x Time	0.9817
SEM	0.575	0.342	0.396		
Cholesterol(mg/egg)					
C	177.83	203.49	190.66	Diet	0.7495
B	180.15	211.49	195.82	Time	0.0016
L	174.01	201.94	187.97	Diet x Time	0.9645
SEM	10.859	6.577	7.446		

53 N= 30eggs pen/group per time (T1 and T2)

54 ^{A,B...} Means within a column lacking a common superscript differ (P<0.05)

55 C: basal diet; L: basal diet + 0.1 % of *Lactobacillus acidophilus*; B: basal diet + 0.05% of *Bacillus*

56 *subtilis*.

57