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## Effects of yellow mealworm larvae ( Tenebrio molitor ) inclusion in diets for female broiler chickens: implications for animal health and gut histology

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1	Effects of yellow mealworm larvae (Tenebrio molitor)
2	inclusion in diets for female broiler chickens: implications
3	for animal health and gut histology
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#### 49 Abstract

50 The aim of the present study was to evaluate the animal 51 performance, haematochemical parameters, intestinal 52 morphology and histological features of broiler chickens fed 53 diets including Tenebrio molitor (TM) larvae meal. A total of 54 160 female broiler chicks (Ross 708) at one-day of age were 55 randomly allotted to four dietary treatments: a control (C) 56 group and three TM groups, in which TM meal was included at 57 50, 100 and 150 g/kg, respectively. Each group consisted of 58 five pens as replicates, with eight chicks per pen. After the 59 evaluation of growth performance and haematochemical 60 parameters, two birds per pen were slaughtered at 40 days and 61 carcass traits were recorded. Morphometric investigations were 62 performed on duodenum, jejunum and ileum and 63 histopathological alterations were assessed for liver, spleen, 64 thymus, bursa of Fabricius, kidney and heart. The live weight 65 (LW) showed a linear (12 days, P < 0.05, maximum with TM15) and quadratic response (40 days, P < 0.05, maximum 66 67 with TM5) to dietary TM meal inclusion. The average daily gain (ADG) showed a linear increase (1-12 days, P < 0.05, 68 69 maximum with TM15) in response to TM meal utilization. A 70 linear effect (1-12 and 12-25 days, P < 0.01 and P < 0.05, 71 maximum with TM15 and TM5) was observed for the daily feed intake (DFI). The feed conversion ratio (FCR) showed a 72 73 linear response to TM utilization in the period 12-25 days (P <

74	0.01, maximum with TM15). A quadratic effect (P $<$ 0.05,
75	maximum with TM5) was observed for the carcass weight. The
76	abdominal fat weight and percentage showed a linear response
77	to dietary TM meal inclusion (P $< 0.05$ and P $< 0.01,$ maximum
78	with TM15 and TM10). A quadratic increase (P $< 0.05$ ,
79	maximum with TM10) was observed for the erythrocytes,
80	while the albumin and GGT showed a linear and quadratic
81	decrease (P < 0.05, minimum with TM10) in relation to TM
82	utilization. Gut morphology and histopathological findings
83	were not significantly influenced (P $> 0.05$ ) by dietary TM
84	meal inclusion. The present study suggests that increasing
85	levels of dietary TM meal inclusion in female broiler chickens
86	diets may improve body weight and feed intake, but can
87	partially worsen feed efficiency. However, positive effects on
88	carcass traits and haematochemical parameters related to TM
89	meal utilization are observed, along with no negative influence
90	on gut morphology and histological findings.

91

# 92 Keywords

93 Poultry; *Tenebrio molitor*; insect meal; growth performance;
94 histology; morphometry.

#### 96 Introduction

97 World population is expected to increase by over a third, 98 reaching over 9 billion people in 2050. This trend suggests that 99 market demand for food will continue to grow. In particular, 100 the demand for cereals and protein sources in both human food 101 and animal feed is projected to have an exponential growth by 102 2050 (FAO, 2013). Consequently, the world supply of some 103 feedstuffs like soybean and maize conventional will 104 increasingly compete between humans and livestock. 105 Therefore, the foremost gamble will be the identification of 106 alternative sources of protein, energy and other nutrients for 107 livestock, in order to avoid such a competition.

108 The potential of insects for becoming a standard ingredient in 109 animal feeds has already been emphasized by several studies 110 (Veldkamp et al., 2012; Van Huis, 2013; Henry et al., 2015), 111 because of the high quality and quantity of protein (Makkar et 112 al., 2014), the low competitiveness with human food (Ballitoc 113 and Sun, 2013) and the reduction of the environmental impact 114 (Oonincx and de Boer, 2012; Makkar et al., 2014; Sánchez-115 Muros et al., 2014). Currently, the considered most valuable 116 insect species to be used in livestock feeds are Hermetia 117 illucens L. (black soldier fly), Musca domestica L. (common 118 house fly), Tenebrio molitor L. (yellow mealworm), Bombyx 119 mori L. (silkworm) and several grasshoppers (Van Huis, 2013). 120 In particular, Tenebrio molitor (TM) and Hermetia illucens

121	have recently been used in poultry (Biasato et al., 2016; Bovera
122	et al., 2016; Schiavone et al., 2017a) and fish (Belforti et al.,
123	2015; Gasco et al., 2016; Renna et al., 2017) feeding. Yellow
124	mealworms are already industrially produced as feed for pets
125	and zoo animals, such as birds, reptiles, small mammals,
126	amphibians and fish, and TM larvae are easily bred on dried
127	waste materials, being able to recycle them into high-quality
128	feed with less energy cost, land area utilization and footprints
129	(Makkar et al., 2014). The influence of TM-based diets on
130	growth performance (Bovera et al., 2015; Bovera et al., 2016;
131	Biasato et al., 2016) haematochemical profile (Bovera et al.,
132	2015; Biasato et al., 2016) and carcass traits (Ballitoc and Sun,
133	2013; Bovera et al., 2016; Bisato et al., 2016), has recently
134	been investigated. Gut morphology, which has been reported to
135	be widely affected by dietary modifications in broilers
136	(Laudadio et al., 2012; Gopinger et al., 2014; Qaisrani et al.,
137	2014), has also been evaluated in free-range chickens fed diets
138	with dietary TM larvae meal inclusion (Biasato et al., 2016).
139	Despite insect meals being considered suitable ingredients for
140	poultry feeding (Veldkamp et al., 2012; van Huis, 2013;
141	Makkar et al., 2014), the implications of their utilization on
142	poultry health and gut development are still very limited. The
143	aim of the present study was to evaluate the growth

144 performance, haematochemical parameters, carcass traits,

intestinal morphology and histological features of femalebroiler chickens fed diets including TM meal.

147

## 148 Materials and Methods

#### 149 Birds and Husbandry

150 The present trial was performed in collaboration with a local poultry corporation named "O.R.A. Agricola S.r.l." sited in 151 152 Cherasco (Cuneo, Italy). The experimental protocol was 153 designed according to the guidelines of the current European 154 and Italian laws on the care and use of experimental animals 155 (European Directive 86 609/EEC, put into law in Italy with 156 D.L. 116/92). Furthermore, the experimental protocol was 157 approved by the Ethical Committee of the Department of 158 Veterinary Sciences of the University of Turin (Italy). A 159 poultry house of 14 m wide  $\times$  141 m long  $\times$  4.7 m high, 160 equipped with waterproof floor and wall, completely covered 161 by tiles and provided with automatic ventilation system was 162 used.

163 A total of 160 female broiler chicks (Ross 708) at one-day of 164 age were randomly allotted to 4 dietary treatments, each 165 consisting of 5 pens as replicates with 8 chicks per pen. Each 166 pen was 1.20 m wide  $\times$  1.20 m long and was equipped with a 167 feeder occupying a surface of almost 1800 cm<sup>2</sup>, three nipple 168 drinkers and rice hulls as litter. During the first three weeks, the 169 animals were heated by infrared lamps to maintain the suitable

170 temperature according to standard breeding practices (Aviagen, 171 2014). Lighting schedule was 23 hours light : 1 hour darkness 172 until day 3 and 18 h light : 6 hours darkness until slaughter age. 173 At hatching, all chicks received vaccination against Newcastle 174 disease, Gumboro disease, infectious bronchitis and 175 coccidiosis. Vaccine recalls were performed on day 9 for 176 infectious bronchitis and on day 18 for Gumboro and 177 Newcastle diseases.

178

#### 179 *Diets*

180 A basal diet based on corn meal, corn gluten meal and soybean 181 meal was formulated and served as control (C) group, while 50, 182 100 and 150 g/kg full-fat TM larvae meal (Gaobeidian 183 Shannong Biology CO., LTD, Gaobeidian, Hebei province, 184 China) inclusion as a partial replacement of soybean meal, corn 185 gluten meal and soybean oil constituted the three experimental 186 treatment groups (TM5, TM10 and TM15) (Table 1). TM meal 187 nutritive composition and energy content were the following: 188 dry matter, 939.0 (g/kg as fed); organic matter, 912.0 (g/kg as 189 fed); crude protein (CP), 519.0 (g/kg as fed); ether extract (EE), 190 236.0 (g/kg as fed); 117.0 (g/kg as fed); neutral detergent fibre 191 (NDF); 79.5 (g/kg as fed); acid detergent fibre (ADF); DL-192 methionine, 10.1 (g/kg as fed); L-lysine, 35.9 (g/kg as fed); 193 gross energy, 24.4; apparent metabolizable energy (AMEn), 194 16.02 (MJ/kg DM). Three different diets were used per each

195 dietary treatments during the three phases of growth: a starter 196 diet (days 1 to 12), a grower diet (days 12 to 25) and a finisher 197 diet (day 25 to 40). For each phase, the experimental diets were 198 isonitrogenous and isoenergetic and were formulated using the 199 AMEn values for TM measured in vivo for broiler chickens (De 200 Marco et al., 2015). Diets met or exceeded NRC (1994) 201 requirements and were adjusted according to Aviagen (2014) 202 broiler nutrition specifications. Feed and water were provided 203 ad libitum.

204

# 205 Chemical analysis

The diets were ground to pass through a 0.5-mm sieve and stored in airtight plastic containers for DM (method number 943.01), ash (method number 924.05), CP (method number 954.01), EE (method number 920.39), NDF (method number 2002.04) and ADF (method number 973.18) determination (AOAC, 2004).

212

## 213 Growth Performances

Health status and mortality were daily monitored during the whole experimental period. Live weight (LW) of the animals was recorded at an individual level at the beginning of the trial, at day 12, 25 and 40. Average daily gain (ADG) and average daily feed intake (DFI) were recorded at an individual and a pen level, respectively, at the end of each growth period. Feed conversion ratio (FCR) was determined for each growth period
and for the overall experimental period. All measurements were
made on the pen basis using a high precision electronic scale
(Sartorius – Signum®).

224

# 225 Slaughtering procedure and recordings

At day 40, all birds were individually weighed and 10
broilers/diet (2 birds/pen) were chosen on the basis of pen
average LW and identified with a shank ring.

229 All selected animals were slaughtered in a commercial abattoir. 230 The plucked and eviscerated carcasses were obtained and the 231 head, neck, feet and abdominal fat were removed to obtain 232 carcass-for-grilling. The weights of liver, spleen, gizzard and 233 abdominal fat were immediately recorded. All slaughtered 234 carcasses were stocked in a cooling chamber (0-4 °C) for 24 h. 235 Weights of carcass-for-grilling, breast and thighs were 236 successively recorded. Carcass-for-grilling, breast, thigh and 237 organs weights were also expressed as percentage of LW.

Collected feet were examined macroscopically using the
Swedish FPD scoring system (Ekstrand et al., 1997). According
to this system, 0 = no lesion, slight discoloration of the skin or
healed lesion; 1= mild lesion, superficial discoloration of the
skin and hyperkeratosis; 2 = severe lesion, affected epidermis,
blood scabs, hemorrhage and severe swelling of the skin.

#### 245 Haematological and serum parameters

246 At slaughter, blood samples were collected from the identified 247 broilers: 2.5 mL was placed in an EDTA tube and 2.5 mL in a 248 serum-separating tube. A blood smear was prepared, using one 249 glass slide for each bird, from a drop of blood without 250 anticoagulant. The smears were stained using May-Grünwald 251 and Giemsa stains (Campbell, 1995). The total red and white 252 blood cell counts were determined in an improved Neubauer 253 haemocytometer on blood samples previously treated with a 254 1:200 Natt-Herrick solution. One hundred leukocytes, including 255 granular (heterophils, eosinophils and basophils) and non-256 granular (lymphocytes and monocytes) leukocytes, were 257 counted on the slide and the heterophils to lymphocytes (H/L) 258 ratio was calculated. The tubes without anticoagulant were left 259 to clot in a standing position at room temperature for 260 approximately two hours to obtain serum. The serum was 261 separated by means of centrifugation at  $700 \times g$  for 15 minutes 262 and frozen at -80°C until analysis. The total proteins were 263 quantified by means of the "biuret method" (Bio Group 264 Medical System kit; Bio Group Medical System, Talamello 265 (RN), Italy); the electrophoretic pattern of the serum was 266 obtained using a semi-automated agarose gel electrophoresis 267 system (Sebia Hydrasys®, Norcross, GA, USA). The alaninoaminotransferase (ALT), aspartate-aminotransferase (AST), 268 269 gamma glutamyl transferase (GGT), triglycerides, cholesterol, glucose, phosphorus, magnesium, iron, uric acid and creatinine
serum concentrations were measured by means of enzymatic
methods in a clinical chemistry analyzer (Screen Master Touch,
Hospitex diagnostics Srl., Firenze, Italy).

274

## 275 Histomorphological investigations

276 The slaughtered animals were submitted to 277 anatomopathological investigations. Intestinal segment samples 278 (approximately 5 cm in length) of duodenum, jejunum, ileum 279 and caecum were excised from each bird and flushed with 0.9% 280 saline to remove all the content. The collected segments of 281 intestine were the loop of the duodenum, the tract before 282 Meckel's diverticulum (jejunum), the tract before the ileocolic 283 junction (ileum) and the apex of the caeca (caecum). Gut 284 segments were fixed in Carnoy's solutions for morphometric 285 analysis. Tissues were routinely embedded in paraffin wax 286 blocks, sectioned at 5 µm thickness, mounted on glass slides 287 and stained with Haematoxylin & Eosin (HE). The evaluated 288 morphometric indices were Vh (from the tip of the villus to the 289 crypt), Cd (from the base of the villus to the submucosa) and 290 the Vh/Cd ratio (Laudadio et al., 2012). Morphometric analyses 291 were performed on 10 well-oriented and intact villi and 10 292 crypts chosen from duodenum, jejunum and ileum (Qaisrani et 293 al., 2014). Samples of liver, spleen, thymus, bursa of Fabricius, 294 kidney and heart were also collected from each animal and

295 fixed in 10% buffered formalin solution for histopathological 296 examination. Tissues were processed in the same way as gut 297 and the following histopathological alterations were evaluated: 298 white pulp hyperplasia and depletion in spleen, cortical 299 depletion in thymus, follicular depletion and intrafollicular 300 cysts in bursa of Fabricius and lymphoid tissue activation in 301 liver (Biasato et al., 2016). Heart and kidney were assessed for 302 inflammatory and degenerative diseases. The observed 303 histopathological findings were evaluated using a 304 semiquantitative scoring system as previously assessed by 305 Biasato et al. (2016): absent/minimal (score = 0), mild (score = 306 1) and severe (score = 2).

307

#### 308 Statistical Analysis

309 IBM SPSS Statistics V20.0.0 software was used to perform 310 statistical analysis. Shapiro-Wilk's test established normality or 311 non-normality of distribution. The experimental unit was the 312 pen for growth performance, haematochemical parameters and 313 carcass traits and bird for histomorphological findings. Data collected for growth performance, blood parameters and 314 315 carcass traits were tested by one-way ANOVA, evaluating the effect of dietary TM inclusion by polynomial contrasts.  $\chi^2$  test 316 317 was performed to evaluate the association between the 318 mortality rate and the dietary treatments. Intestinal 319 morphometric indices were analyzed by fitting a general linear

320 model (GLM). The GLM allowed the morphometric indices 321 (Vh, Cd and Vh/Cd, separately) to depend on three fixed 322 factors (diet, intestinal segment and interaction between diet 323 and intestinal segment). The interactions between the levels of 324 the fixed factors were evaluated by pairwise comparisons. 325 Statistical analysis was performed by procedure "General 326 Linear Models > Univariate". Histopathological and FPD 327 scores were analyzed by Kruskal-Wallis test (post-hoc test: 328 Dunn's Multiple Comparison test). P values < 0.05 were 329 considered statistically significant. The results were expressed 330 as mean and pooled standard error of the mean (SEM).

331

# 332 **Results**

#### 333 Growth performance

334 No clinical signs were observed and the birds remained healthy 335 during the whole experimental period. The mortality rates of C 336 (2.5%), TM5 (2.5%), TM10 (0%) and TM15 (2.5%) groups 337 were not significantly different among treatments (P > 0.05). 338 Growth performance of the broiler chickens are summarized in 339 Table 2. At 12 days of age, the LW increased linearly with 340 increasing TM meal levels (P < 0.05) and the linear response 341 increased to a maximum corresponding to the inclusion of 150 342 g/kg of TM meal. At 25 days of age, no significant effects related to TM meal utilization were observed. At 40 days of 343 344 age, the LW showed quadratic response to increasing TM meal

345	levels (P < 0.05), with a maximum corresponding to the
346	inclusion of 50 g/kg of TM meal. In the period from 1-12 days
347	of age, the ADG increased linearly with increasing TM meal
348	levels (P < 0.05) and the linear response increased to a
349	maximum corresponding to the inclusion of 150 g/kg of TM
350	meal. On the contrary, the ADG showed no differences (P $\!$
351	0.05) in the periods from 12 to 25 and 25 to 40 days of age In
352	the period from 1 to 12 days of age, the response of DFI to the
353	effect of TM meal inclusion was statistically significant (P <
354	0.01). In particular, the linear response increased to a maximum
355	corresponding to the inclusion of 150 g/kg of TM meal. In the
356	period from 12 to 25 days of age, the DFI increased linearly
357	with increasing TM meal levels (P $<$ 0.05) and the linear
358	response increased to a maximum corresponding to the
359	inclusion of 50 g/kg of TM meal. On the contrary, the DFI
360	showed no differences (P > 0.05) in the period from 25 to 40
361	days of age. In the periods from 1 to 12, 25 to 40 and 1 to 40
362	days of age, the FCR was similar among the dietary treatments
363	(P > 0.05). Differently, the FCR showed linear response to
364	increasing TM meal levels in the period from 12 to 25 (P $<$
365	0.01), with a maximum corresponding to the inclusion of 150
366	g/kg of TM meal.
267	

368 Slaughtering performance and footpad dermatitis (FPD)
369 score

370 Table 3 summarizes the slaughtering performance of the broiler 371 chickens. The carcass weight increased quadratically with 372 increasing TM meal levels (P < 0.05) and the quadratic 373 response increased to a maximum corresponding to the 374 inclusion of 50 g/kg of TM meal. The abdominal fat weight 375 showed linear response to increasing TM meal levels (P < 376 0.05), with a maximum corresponding to the inclusion of 150 377 g/kg of TM meal. Similarly, the abdominal fat percentage 378 increased linearly with increasing TM meal levels (P < 0.01) 379 and the linear response increased to a maximum corresponding 380 to the inclusion of 100 g/kg of TM meal. On the contrary, no 381 significant effects related to TM meal utilization were observed 382 for the other carcass traits (P > 0.05). FPD scores (C: 0.40; 383 TM5: 0.20; TM10: 0.20; TM15: 0.00) were also not influenced 384 by dietary TM meal inclusion (P > 0.05).

385

# 386 Haematological and serum parameters

387 Haematological and serum biochemical traits of the broiler 388 chickens are summarized in Table 4. The erythrocytes 389 increased quadratically with increasing TM meal levels (P <390 (0.05) and the quadratic response increased to a maximum 391 corresponding to the inclusion of 100 g/kg of TM meal. The 392 albumin showed linear response to increasing TM meal levels (P < 0.05), with a minimum corresponding to the inclusion of 393 100 g/kg of TM meal. Similarly, the GGT decreased 394

395 quadratically with increasing TM meal levels (P < 0.05) and the 396 quadratic response decreased to a minimum corresponding to 397 the inclusion of 100 g/kg of TM meal. On the contrary, no 398 significant effects related to TM meal utilization were observed 399 for the other blood and serum parameters (P > 0.05).

400

#### 401 *Histomorphological investigations*

402 The effects of the diet, intestinal segment and interaction 403 between diet and intestinal segment on gut morphometric 404 indices of the broiler chickens are shown in Tables 5 and 6. 405 There was no influence of diet or interaction between diet and 406 intestinal segment (P > 0.05) on the gut morphometric indices. 407 On the contrary, Vh, Cd and Vh/Cd depended on intestinal 408 segment (P < 0.001, P = 0.001 and P < 0.001, respectively) 409 (Table 5). In particular, the duodenum showed higher Vh (P <410 0.001) than the jejunum and ileum. Furthermore, higher Cd (P 411 < 0.001) was found in the duodenum and jejunum than the 412 ileum. Similarly, the duodenum and jejunum showed higher 413 Vh/Cd (P < 0.001) than the ileum (Table 6).

414 Histopathological alterations were observed in all the dietary 415 treatments and developed in spleen, thymus, bursa of Fabricius 416 and liver, while heart and kidney showed no significant 417 findings. Spleen (C =  $1.50 \pm 0.22$ ; TM5 =  $1.20 \pm 0.25$ ; TM10 = 418  $1.10 \pm 0.31$ ; TM15 =  $1.10 \pm 0.28$ ), thymus (C =  $0.10 \pm 0.10$ ; 419 TM5 =  $0.30 \pm 0.15$ ; TM10 and TM15 =  $0.10 \pm 0.10$ ), bursa of

420	Fabricius (C = 1.90 $\pm$ 0.10; TM5 = 1.50 $\pm$ 0.27; TM10 = 1.80 $\pm$
421	0.20; TM15 = 1.70 $\pm$ 0.21) and liver (C = 0.90 $\pm$ 0.23; TM5 =
422	$1.10 \pm 0.23$ ; TM10 = 0.90 ± 0.23; TM15 = 0.30 ± 0.15) scores
423	were not influenced by dietary TM meal inclusion. Spleen
424	showed moderate (C = $30\%$ of the broilers; TM5 = $40\%$ ; TM10
425	= 10%; TM15 = 30%) to severe (C = $60\%$ ; TM5 = $40\%$ ; TM10
426	= 50%; TM15 = 40%) white pulp depletion or hyperplasia. The
427	10% I, 20% (TM5), 40% (TM10) and 30% (TM15) of the
428	animals had normal spleen (Fig. 1A-B). In thymus, moderate
429	cortical depletion was found in all the groups ( $C = 10\%$ of the
430	broilers; $TM5 = 30\%$ ; $TM10$ and $TM15 = 10\%$ ). A normal
431	thymus was observed in 90% I, 70% (TM5) and 90% (TM10
432	and TM15) of the animals. Bursa of Fabricius showed moderate
433	(C and TM5 = 10% of the broilers; $TM10 = 0\%$ ; $TM15 = 10\%$ )
434	to severe (C = $80\%$ ; TM5 = $70\%$ ; TM10 = $90\%$ ; TM15 = $80\%$ )
435	follicular depletion. The 0% I, 20% (TM5), 10% (TM10) and
436	10% (TM15) of the animals had normal bursa of Fabricius (Fig.
437	1C-D). In liver, moderate (C, TM5 and TM10 = $50\%$ of the
438	broilers; TM15 = 30%) to severe (C = 20%; TM5 = 30%;
439	TM10 = 20%; TM15 = 0%) perivascular lymphoid tissue
440	activation. A normal liver was observed in 30% I, 20% (TM5),
441	30% (TM10) and 70% (TM15) of the animals.

# **Discussion**

# *Growth performances*

Growth performances of the broiler chickens of the present
study were consistent with the reference values recorded in the
commercial farm in which the trial was conducted.

448 The body weight, weight gain and feed intake of the birds in 449 the present trial improved with increasing levels of TM meal 450 inclusion, but the feed efficiency resulted partially impaired. 451 Little information on the influence of dietary TM meal 452 inclusion in broiler chickens is currently available. Ramos-453 Eldoury et al. (2002) and Biasato et al. (2016) did not show any 454 effects for the growth performance in fast-growing and 455 intermediate-growing chickens, respectively, fed diets in which 456 the TM inclusion level ranged from 50 to 100 g/kg. Differently, 457 Ballitoc and Sun (2013) and Bovera et al. (2015) observed 458 improved growth performance in fast-growing chickens fed 459 diets with low (from 5 to 100 g/kg) or high (296 g/kg) TM inclusion levels, respectively. Similar findings were obtained 460 461 by Loponte et al. (2017) in partridges fed diets in which the TM 462 inclusion level ranged from 250 to 500 g/kg. Recently, Islam 463 and Yang (2017) also found a positive effect of a mealworm-464 based probiotic on broiler growth performance. Some authors 465 also explored the possibility to use other insect meals in poultry 466 feeding. Adeniji (2007) and Hwangbo et al. (2009) studied the 467 effects of housefly-maggots as feed supplement in the diet of 468 broiler chickens: the first found no differences for growth 469 performance with inclusion levels ranging from 55 to 220 g/kg, 470 while the latter (inclusion rate: 50-200 g/kg) observed a linear 471 increase in LW gain. Ijaiya and Eko (2009) evaluated the 472 effects of replacing dietary fishmeal with silkworm meal 473 (inclusion rate: 22-93 g/kg) on growth performance of broiler 474 chickens, finding no differences related to insect meal 475 utilization. Also Oyegoke et al. (2006) and Wang et al. (2005) 476 observed no adverse effects on growth performance of broiler 477 chickens fed diets with Cirina forda (inclusion rate: 20-40 478 g/kg) and Gryllus testaceus (inclusion rate: 50-150 k/kg), 479 respectively. Cullere et al. (2016) recently studied the influence 480 of the inclusion (from 100 to 150 g/kg) of Hermetia illucens 481 meal in quail diet, finding no differences for growth 482 performances. Schiavone et al. (2017b) also evaluated the 483 effects of replacing soybean oil with Hermetia illucens meal 484 (inclusion rate: 500-1000 g/kg) on growth performance of 485 broiler chickens, finding no differences related to insect meal 486 utilization. The wide variability of the results obtained in the 487 previous studies may be related to the nutritive value of the 488 insect meal used, which can be influenced by the species, the 489 insect life stage (adult, larva or pupa) and the insect rearing 490 substrate (Sànchez-Muros et al., 2014).

491 The improvement of feed intake observed in the birds fed TM 492 diets of the present trial was considered suggestive of increased 493 feed palatability in relation to the addition of yellow 494 mealworms, since insects are naturally consumed by wild birds

495 and free-range poultry (Zuidhof et al., 2003). In particular, the 496 increased DFI observed in the starter period, which was 497 accompanied by increased LW and ADG and unaffected FCR, 498 was quite relevant. Indeed, starter period (from hatch to 10 499 days) is considered the most important in broiler production, 500 since growth and development take place at an incredible rate 501 during it. In this period, the chicks' weight quadruples, thus 502 influencing the following growth rate (Aviagen, 2014). On the 503 contrary, the increased DFI observed in the growing period was 504 accompanied by unaffected LW and ADG and subsequently 505 impaired FCR, thus representing a negative effect related to 506 TM meal utilization. De Marco et al. (2015) speculated that the 507 chitin contained in the exoskeleton of the TM meal may 508 negatively influence the apparent digestibility coefficient of the 509 total tract of nutrients. Furthermore, Ravindran and Blair 510 (1993) pointed out that the chitin of insects is difficult to digest 511 by domestic poultry. As suggested by Rumpbold and Schlüter 512 (2013), the partial chitin removal through high pressure 513 processing could improve the use of insects as feeding 514 ingredient thanks to disruption of the link between some chitin-515 bound proteins. However, the limited number of birds included 516 in the current trial could have influenced the data interpretation. 517 The results obtained need to be confirmed on a larger number 518 of animals.

519

# 520 Slaughtering performance and footpad dermatitis (FPD) 521 score

522 The majority of the carcass traits of the broilers in the present 523 trial were not influenced by dietary TM meal inclusion, as 524 previously observed by Bovera et al. (2016) and Biasato et al. 525 (2016). Similar findings were obtained by Cullere et al. (2016) 526 and Schiavone et al. (2017) in broiler quails and chickens fed 527 diets with Hermetia illucens meal and fat, respectively. 528 However, the carcass weight, abdominal fat weight and 529 abdominal fat percentage increased with increasing levels of 530 TM meal utilization. Loponte et al. (2017) also observed 531 improved carcass weights when TM and Hermetia illucens 532 meals were included in the diets of partridges. As already 533 suggested by them, the differences in the eviscerated carcass 534 weights can be partially explained by the increased final LW of 535 the birds. Similar findings in terms of improved eviscerated 536 carcass weights were obtained by Khatun et al. (2003), 537 Hwangbo et al. (2009) and Ballitoc and Sun (2013), who also 538 observed improved slaughter, dressed carcass, breast muscle 539 and thigh muscle weights and dressing percentage in broilers 540 fed diets with different insect meals inclusion. The differences 541 in the abdominal fat weight and abdominal fat percentage 542 observed in the birds of the present study are also in agreement with Ballitoc and Sun (2013) and suggests that yellow 543

544 mealworm utilization may improve fat mass in broiler chickens545 (USDA, 2011).

The majority of FPD scores obtained in the present trial was zero and no differences were found in relation to TM meal utilization. This is a positive result, since a low prevalence and severity of FPD is highly desirable as far as health of birds and product quality are concerned (Meluzzi et al., 2008).

551

# 552 Haematological and serum parameters

553 All the blood parameters obtained in the present trial fell within 554 the physiological ranges (Lumej, 2008), thus suggesting that 555 TM meal utilization does not affect health status of the animals. 556 In particular the H/L ratio, that is commonly used as indicator 557 of stress in poultry (De Marco et al., 2013; Salamano et al., 558 2010), was not affected by dietary TM meal inclusion. As 559 already observed by Bovera et al. (2015) and Biasato et al. 560 (2016), the majority of the haematochemical and serum 561 biochemical traits were not affected by yellow mealworm 562 inclusion in the birds of the present study. Similar findings 563 were obtained by Schiavone et al. (2017b) in broilers fed diets 564 with Hermetia illucens fat. However, the erythrocytes increased 565 and albumin and GGT decreased with increasing levels of TM 566 meal utilization Loponte et al. (2017) also observed lower albumin when TM and Hermetia illucens meals were included 567 in the diets of partridges. Interestingly, this finding was 568

569 accompanied by the increase of albumin/globulin ratio, which 570 was also reported by Bovera et al. (2015) and ascribed to the 571 properties of chitin contained in insect meal. High globulin 572 concentrations and low albumin/globulin ratios generally 573 indicate better disease resistance and immune response of birds 574 (Griminger and Scanes, 1986). Another interesting finding is 575 represented by the decrease of GGT serum levels in TM 576 animals. Indeed, a high GGT concentration in birds is used as 577 an indicator of liver disease and bile flow disorders (Ognik and 578 Krauze, 2016). Therefore, GGT reduction can be considered a 579 positive effect related to TM meal utilization.

580

#### 581 Histomorphological investigations

582 Dietary TM meal inclusion did not influence the gut 583 morphology of the birds of the present study, as already 584 observed by Biasato et al. (2016). Morphometric measurements 585 of Vh and Cd are generally used to assess intestinal 586 development (Franco et al., 2006), since represent useful 587 indicators of gut proliferative and absorptive compartments 588 (Lenhardt and Mozes, 2003). The Vh/Cd ratio is also evaluated, 589 because it gives an indication of the likely maturity and 590 functional capacity of the enterocytes (Hampson, 1986). As 591 previously reported (Uni et al., 1999; Iji et al., 2001; Biasato et 592 al., 2016), the present study confirms that both duodenum and 593 jejunum show a greater morphological development compared 594 with the ileum. Indeed, the duodenum is the intestinal tract with 595 the fastest cell renewal, and is also the first segment of the 596 small intestine to receive physical, chemical and hormonal 597 stimuli provoked by diet (Macari, 1998). Furthermore, the 598 jejunum is an important site for nutrient digestion (Iji et al., 599 Therefore, dietary TM inclusion 2001). preserves a 600 proximodistal decreasing gradient of the morphometric indexes 601 from the duodenum to the ileum, thus suggesting the 602 maintenance of the physiological gut development.

603 The broiler chickens of the present trial showed different 604 degrees of lymphoid system activation, with no differences 605 related to dietary TM meal inclusion. This result could be 606 related to the stress occurrence in modern poultry rearing 607 operations. Stress can be caused by a variety factors, 608 physiological (rapid growth rate) and social (overcrowding) 609 ones included (Liles et al., 2015). However, a great deal of 610 individual bird variability in some immunological measures 611 (i.e., stimulation index, heterophils to lymphocyte ratio and 612 lymphocyte blastogenesis) may also be considered (Talebi et 613 al., 1995).

614

#### 615 Conclusion

In conclusion, the present study suggests that increasing levels
of dietary TM meal inclusion in female broiler chickens diets
may improve body weight, weight gain and feed intake, but can

619 partially worsen feed efficiency. However, positive effects on 620 carcass traits and haematochemical parameters are observed, 621 along with no negative influence on gut morphology and 622 histological findings. These results confirm previous data 623 concerning the safety of TM utilization in poultry feed, even if 624 legislative issues are still needed to allow insect meal to be 625 used as transformed animal protein to feed monogastric farm 626 animals.

627

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860	Table 1. Ingredients (g/kg as fed)	, apparent metabolizable energy	(MJ/kg DM) and nutrient	t composition (%) of the
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## 861 experimental diets.

Incredients	First P	eriod (d	ays 1 to	12)	Second p	eriod (d	ays 12 to	o 25)	Third perio	d (day 2	5 to sla	ughter)
Ingredients	Control	TM5	TM10	TM15	Control	TM5	TM10	TM15	Control	TM5	TM10	TM15
Corn meal	483.2	482.7	488.5	496.6	523.8	535.9	549.3	566.8	566.6	572.2	585.7	605.4
Soybean meal	345.0	333.8	304.0	262.0	317.0	294.0	254.1	203.9	275.5	259.0	219.0	164.0
TM meal	0.0	50.0	100.0	150.0	0.0	50.0	100.0	150.0	0.0	50.0	100.0	150.0
Corn gluten meal	75.5	42.0	23.0	14.5	58.3	24.0	8.0	0.0	56.0	21.0	5.0	0.0
Soybean oil	54.0	50.3	43.5	34.8	64.9	59.9	51.7	41.6	68.9	64.9	56.7	45.8
Dicalcium phosphate	11.0	12.0	13.0	15.5	8.4	9.0	10.5	12.9	7.0	8.0	9.5	12.0
Calcium carbonate	17.5	16.5	16.0	15.0	15.0	15.0	14.5	13.0	14.5	14.0	13.5	12.2
Sodium chloride	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3
Sodium bicarbonate	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3
DL-methionine	0.8	0.9	0.9	0.8	0.8	0.9	0.9	0.8	0.4	0.6	0.6	0.5
L-lysine	3.1	1.9	1.2	0.9	2.0	1.3	0.9	0.8	1.3	0.4	0.0	0.0
Threonine	0.1	0.1	0.1	0.1	0.0	0.2	0.3	0.4	0.0	0.1	0.2	0.3
Trace mineral-vitamin premix <sup>1</sup>	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0
Coline	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2
3-phytase	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Total	100	100	100	100	100	100	100	100	100	100	100	100
AMEn <sup>2</sup> (MJ/kg)	12.89	12.89	12.89	12.89	13.28	13.28	13.28	13.28	13.54	13.54	13.54	13.54
Nutrient composition (%)												
DM	86.6	86.6	86.7	86.6	86.7	86.8	86.6	86.8	86.8	86.7	86.7	86.8
СР	23.5	23.5	23.6	23.8	21.3	21.1	21.1	21.1	19.6	19.6	19.6	19.6
EE	7.9	8.3	9.0	9.6	9.0	9.2	9.8	10.3	9.5	9.7	10.4	10.8
NDF	9.4	9.8	10.0	10.1	9.4	9.8	10.0	10.1	9.4	9.7	9.9	10.1
ADF	3.8	4.1	4.3	4.4	3.7	3.9	4.1	4.1	3.5	3.8	3.9	4.0

Nutrient composition $(\%)^2$												
Calcium	1.1	1.1	1.1	1.1	0.9	0.9	0.9	0.9	0.8	0.8	0.8	0.8
Available phosphorus	0.6	0.6	0.5	0.6	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
Digestible methionine	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.4	0.4	0.4	0.4
Digestible lysine	1.4	1.4	1.4	1.4	1.3	1.3	1.3	1.3	1.1	1.1	1.1	1.1
Digestible threonine	0.9	0.9	0.9	0.9	0.9	0.9	0.9	0.9	0.8	0.8	0.8	0.8

<sup>862 &</sup>lt;sup>1</sup>Mineral-vitamin premix (Final B Prisma, IZA SRL), given values are supplied per kg of diet: 2.500.000 IU of vitamin A;

864 mg of vitamin B6; 4 mg of vitamin B12; 30 mg of biotin; 3.111 mg of Ca pantothenate acid; 100 mg of folic acid; 15.000 mg

865 of vitamin C; 5.600 mg of vitamin B3; 10.500 mg of Zn, 10.920 mg of Fe; 9.960 mg of Mn; 3.850 mg of Cu; 137 mg of I; 70

866 mg of Se.

<sup>2</sup>Calculated according to INRA 2004 and De Marco et al. (2015).

868 TM, Tenebrio molitor; AME, apparent metabolizable energy; DM, dry matter; CP, crude protein; EE, ether extract; NDF,

869 neuter detergent fiber; ADF, acid detergent fiber.

<sup>863 1.000.000</sup> IU of vitamin D3; 7.000 IU of vitamin E; 700 mg of vitamin K; 400 mg of vitamin B1; 800 mg of vitamin B2; 400

			Dietary	treatments <sup>1</sup>				$\mathbf{P}^3$
Variable <sup>2</sup>	Age	С	TM5	TM10	TM15	SEM	Linear	Quadratic
	12 d	303.15	338.25	339.15	351.83	6.78	0.013	0.352
LW (g)	25 d	1174.90	1234.77	1183.35	1179.79	14.14	0.775	0.280
	40 d	2078.46	2309.97	2115.91	2084.22	30.51	0.408	0.012
	1-12 d	23.61	26.80	26.89	28.04	0.61	0.012	0.350
ADG (g)	12-25 d	67.06	68.96	64.94	63.69	1.17	0.198	0.512
	25-40 d	60.24	71.68	62.17	60.29	1.91	0.552	0.071
	1-12 d	25.40	28.80	30.80	32.40	0.95	0.006	0.583
DFI (g)	12-25 d	95.00	117.00	109.40	116.60	2.95	0.014	0.129
	25-40 d	184.41	220.23	209.49	225.28	6.85	0.731	0.741
	1-12 d	1.08	1.07	1.14	1.15	0.03	0.336	0.925
$\mathbf{ECD}\left( \mathbf{r}\left( \mathbf{r}\right) \right)$	12-25 d	1.42	1.74	1.68	1.86	0.05	0.001	0.325
FCR (g/g)	25-40 d	2.40	2.16	2.13	2.31	0.11	0.775	0.404
	1-40 d	1.78	1.84	1.81	1.95	0.05	0.342	0.730

871 **Table 2.** Effect of the dietary TM larvae meal inclusion on the growth performance of the female broiler chickens.

872 Each mean represents 5 replicates with 8 chicks/replicate (n = 40/treatment).

 $^{1}$ Four dietary treatments: C = control; TM5 = 5% inclusion level of *Tenebrio molitor*; TM10 = 10% inclusion level

- 874 of *Tenebrio molitor*; TM15 = 15% inclusion level of *Tenebrio molitor*.
- <sup>2</sup>LW, live weight; DFI, daily feed intake; ADG, average daily gain; FCR, feed conversion ratio; n, number of pens.

876 <sup>3</sup>Statistical significance: P < 0.05.

		Dietar	ry treatments <sup>2</sup>		$\mathbf{P}^3$		
Variable	С	TM5	TM10	TM15	SEM	Linear	Quadratic
Live weight (LW) (g)	1980	2149	2118	2018	33.68	0.771	0.054
Carcass weight (g)	1377	1501	1500	1276	38.80	0.348	0.025
Carcass weight (% LW)	69.46	69.75	70.77	62.86	1.78	0.256	0.266
Breast (g)	332	388	359	353	10.24	0.688	0.135
Breast (% LW)	16.64	17.92	16.94	17.43	0.24	0.512	0.406
Thigh (g)	411	433	434	402	6.00	0.646	0.026
Thigh (% LW)	20.80	20.20	20.51	20.00	0.16	0.163	0.883
Spleen (g)	2.62	2.84	2.83	2.80	0.13	0.672	0.677
Spleen (% LW)	0.13	0.13	0.13	0.14	0.01	0.742	0.768
Liver (g)	25.01	27.30	27.03	25.68	0.82	0.825	0.302
Liver (% LW)	1.26	1.26	1.27	1.26	0.02	0.921	0.941
Gizzard (g)	36.75	38.20	36.83	34.40	1.03	0.392	0.379
Gizzard (% LW)	1.83	1.77	1.74	1.69	0.04	0.168	0.914
Abdominal fat (g)	40.5	43.3	33.8	43.5	1.29	0.014	0.057
Abdominal fat (% LW)	0.66	0.86	1.08	0.98	0.23	0.005	0.074

878 **Table 3.** Effect of the dietary TM larvae meal inclusion on the carcass traits of the female broiler chickens.<sup>1</sup>

<sup>1</sup>Each mean represents 5 pens with 2 chicks/pen (n = 5/treatment).

880 <sup>2</sup>Four dietary treatments: C = control; TM5 = 5% inclusion level of *Tenebrio molitor*; TM10 = 10% inclusion level

881 of *Tenebrio molitor*; TM15 = 15% inclusion level of *Tenebrio molitor*.

882 <sup>3</sup>Statistical significance: P < 0.05.

		Dietar	y treatments <sup>2</sup>		$\mathbf{P}^4$		
Variable <sup>3</sup>	С	TM5	TM10	TM15	SEM	Linear	Quadratic
Erythrocyte $(10^6 \text{ cell/}\mu\text{l})$	2.27	2.41	2.47	2.39	0.28	0.073	0.040
Leukocyte ( $10^3$ cell/µl)	9.28	9.07	9.31	9.70	0.23	0.521	0.562
H/L ratio	0.83	0.68	0.75	0.77	0.03	0.632	0.095
Albumin (g/dl)	1.66	1.35	1.27	1.32	0.06	0.046	0.134
Total protein (g/dl)	3.31	3.80	3.85	4.10	0.14	0.068	0.663
GGT (UI/l)	26.86	22.13	21.61	25.56	1.05	0.629	0.046
AST (UI/l)	189.48	227.18	217.11	211.06	8.62	0.494	0.229
ALT (UI/l)	20.06	21.33	18.57	15.90	1.16	0.164	0.428
Uric Acid (mg/dl)	3.44	4.03	3.28	3.35	0.21	0.617	0.562
Creatinine (mg/dl)	0.36	0.38	0.36	0.38	0.00	0.410	0.957
Triglycerides (mg/dl)	42.44	44.48	52.80	36.03	3.05	0.686	0.133
Cholesterol (mg/dl)	60.33	71.12	70.87	77.80	3.71	0.138	0.799
Glucose (mg/dl)	222.60	221.10	219.70	227.00	1.62	0.431	0.196
Phosphorus (mg/dl)	3.67	4.19	3.93	5.56	0.33	0.067	0.376
Magnesium (mEq/l)	1.30	1.15	1.18	1.15	0.03	0.159	0.380
Iron (µg/dl)	102.44	81.20	81.26	103.89	11.64	0.985	0.392

## **Table 4.** Effect of the dietary TM larvae meal inclusion on the haematological and serum parameters of the female

885 broiler chickens.<sup>1</sup>

886 <sup>1</sup>Each mean represents 5 pens with 2 chicks/pen (n = 5/treatment).

- 887 <sup>2</sup>Four dietary treatments: C = control; TM5 = 5% inclusion level of *Tenebrio molitor*; TM10 = 10% inclusion level
- 888 of *Tenebrio molitor*; TM15 = 15% inclusion level of *Tenebrio molitor*.
- <sup>3</sup>H/L, heterophiles to lymphocytes ratio; GGT, gamma glutamyl transferase; AST, aspartate aminotransferase;
- 890 ALT, alanine aminotransferase; n, number of birds.
- 891 <sup>4</sup>Statistical significance: P < 0.05.
- 892

894 **Table 5.** Effects of diet, intestinal segment and interaction between diet and intestinal segment on the intestinal

Index	Fixed effect	d.f. <sup>3</sup>	F	<b>P</b> <sup>4</sup>
	Diet <sup>1</sup>	3	1.210	0.310
Vh (mm)	Intestinal segment <sup>2</sup>	2	68.115	< 0.001
	$\text{Diet} \times \text{Intestinal segment}$	6	0.922	0.483
	Diet	3	0.891	0.449
Cd (mm)	Intestinal segment	2	7.275	0.001
	$\text{Diet} \times \text{Intestinal segment}$	6	1.593	0.157
	Diet	3	0.705	0.551
Vh/Cd (mm/mm)	Intestinal segment	2	35.195	< 0.001
	$Diet \times Intestinal segment$	6	0.277	0.947

895 morphometric indices of the female broiler chickens.

- 896 <sup>1</sup>Four dietary treatments: C = control; TM5 = 5% inclusion level of *Tenebrio molitor*; TM10 = 10% inclusion level
- 897 of *Tenebrio molitor*; TM15 = 15% inclusion level of *Tenebrio molitor*.
- 898 <sup>2</sup>Three intestinal segments: duodenum, jejunum and ileum.
- <sup>3</sup>Degrees of freedom.
- 900 <sup>4</sup>Statistical significance: P < 0.05.

901 Vh, villus height; Cd, crypt depth; Vh/Cd, villus height to crypt depth ratio.

902	Table 6. Least square me	ans of intestinal morphometric	c indices in female broiler chickens in
	1	1	

Index	Fixed effect	Effect levels	Least square mean <sup>1</sup>	SEM	
		С	1.73		
	Diet <sup>2</sup>	TM5	1.67	0.06	
	Diet	TM10	1.57	0.06	
Vh (mm)		TM15	1.61		
		DU	2.08 <sup>a</sup>		
	Intestinal segment <sup>3</sup>	JE	1.65 <sup>b</sup>	0.05	
		Ι	1.20 <sup>c</sup>		
		С	0.20		
	Diet	TM5	0.20	0.01	
		TM10	0.20		
Cd (mm)		TM15	0.21		
		DU	0.21 <sup>a</sup>		
	Intestinal segment	JE	0.21 <sup>a</sup>	0.00	
		Ι	0.19 <sup>b</sup>		
		С	8.49		
	Diet	TM5	8.51	0.34	
	Diet	TM10	8.00	0.54	
Vh/Cd (mm/mm)		TM15	8.00		
		DU	10.06 <sup>a</sup>		
	Intestinal segment	JE	8.13 <sup>b</sup>	0.47	
		Ι	6.56 <sup>c</sup>		

903 relation to diet and intestinal segment.

904 <sup>1</sup>Means with different superscript letters (a, b) within the same column per fixed effect (i.e.

905 diet, intestinal segment) differ significantly (P < 0.05).

- $^{2}$ C = control; TM5 = 5% inclusion level of *Tenebrio molitor*; TM10 = 10% inclusion level of
- *Tenebrio molitor*; TM15 = 15% inclusion level of *Tenebrio molitor*
- $^{3}$ DU = duodenum; JE = jejunum; I = ileum.

## 909 Figure legends

Figure 1. Histological findings of the female broiler chickens. A) TM5 group. A normal
spleen. 5× Haematoxylin & Eosin stain. B) TM5 group. Spleen with severe and diffuse
depletion of the white pulp. A high number of apoptotic cells (arrowheads) are observed. 20×
Haematoxylin & Eosin stain. C) C group. A normal follicle in the bursa of Fabricius. 10×
Haematoxylin & Eosin stain. D) C group. Bursa of Fabricius with mild and multifocal
follicular depletion (arrow) associated with intrafollicular cyst (\*). 10× Haematoxylin &