

1 **Growth performance, clinical evaluation and sensory impact of black soldier fly larval**  
2 **meal as protein resource on grower-finisher guinea fowls reared under tropical**  
3 **conditions**

4 <sup>1\*</sup>Wallace, P. A., <sup>1</sup>Nyameasem, J. K., <sup>1</sup>Aboagye, G. A., <sup>1</sup>Affedzie-Obresi, S., <sup>1</sup>Nkegbe, K.,  
5 <sup>3</sup>Murray, F., Botchway, V., <sup>1</sup>Karbo, N., <sup>3</sup>Leschen, W., <sup>3</sup>Maquart, P-O. and <sup>2</sup>Clottey, V.

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7 <sup>1</sup>Council for Scientific and Industrial Research - Animal Research Institute, P. O. Box AH 20,  
8 Achimota, Accra, Ghana

9 <sup>2</sup>CABI-WAC, P. O. Box 860, Cantoments, Accra, Ghana

10 <sup>3</sup>University of Stirling, Stirling, FK9 4LA, United Kingdom

11

12 **Corresponding author:**

13 CSIR-Animal Research Institute, P. O. Box AH 20, Achimota, Accra, Ghana

14 E-mail: [pwallaus@yahoo.com](mailto:pwallaus@yahoo.com)

15 Tel.: +233-50-4072177

16

17 **Abstract**

18 The study was conducted with the view to determine the impact that larval meal from black  
19 soldier fly (BSFLM) would have on growing guinea fowls when used utilized as fishmeal  
20 replacer. BSFLM, produced from decaying mango fruits, were harvested, dried, milled and  
21 used for the feeding trial. BSFLM replaced fish meal in the ratios of 0, 20, 40, 60, 80 and  
22 100% to produce six dietary treatments which were iso-caloric and iso-nitrogenous. Two  
23 hundred and forty eight-week old grower guinea fowls with mean live-weight of  $273.2 \pm 10.9$   
24 g were tagged, weighted and randomly assigned to six floor pens. Each bird was treated as a  
25 replicate. Feed and water were provided *ad libitum*. During the entire period which lasted ten  
26 weeks. Feed consumption differed among the treatment groups ( $P = 0.0072$ ) with the 100%  
27 fishmeal diets recording the lowest. However, daily gain was found to be significantly  
28 ( $P=0.009$ ) higher for birds fed high BSFL diets compared to the control (fishmeal diet). The  
29 inclusion of BSFLM in the diets seemed to have elicited positive linear effect on the weight  
30 gains of the guinea fowls ( $R^2 = 0.91$ ) with increasing concentration resulting in higher live  
31 weight gains. The FCR also differed between treatments ( $P<0.05$ ) but similar for 100%  
32 fishmeal (control) and 100 % BSFLM diets. The study further revealed that BSFLM  
33 replacement of fish meal in guinea fowl diets would not adversely affect the haematopoietic  
34 ability of the birds. Organ and haematopoietic integrity were equally assured regardless of the  
35 protein types used as well as levels of inclusion.

36 **Keywords:** blood chemistry, carcass, haematology, organoleptic properties, protein

37

## 38 **Introduction**

39 Poultry production has been advanced to provide a rapid means of producing animal protein  
40 to meet the nutritional requirements of the ever increasing human populace (Taiwo et al.  
41 2005). However, sustainable feeding of poultry has been a major setback in sub-Saharan  
42 African resulting in huge imports of both poultry products and feedstuffs. The search for safe  
43 and suitable but equally nutritious protein alternatives to meet the well-known challenge of  
44 high cost of feed resulting from soaring prices of the conventional feedstuffs such as fishmeal  
45 and soybean (Mmereole 2008; Dei et al. 2013) continue unabated. Coupled to this, is the  
46 issue of their seasonal unavailability and stiff competition with humans who usually use them  
47 as part of the staple meal. Additionally, the increasing human population has been reported to  
48 require a corresponding increasing demand for the conventional feed ingredients (Dei et al.  
49 2013).

50 The black soldier fly (*Hermetia illucens* Linnaeus 1758) is a fly (Diptera) of the  
51 *Stratiomyidae* family (Tran et al. 2015) which has been proposed since the 1990s and actually  
52 tested as an efficient way of disposing of organic waste into fat- and protein-rich biomass  
53 suitable for various beneficial applications such as protein source for all livestock and poultry  
54 among others (Diener et al. 2011; van Huis et al. 2013; Wallace et al. 2017). Moreover, the  
55 major advantage that *Hermetia illucens* has over other insect species used for biomass  
56 production is that the adult does not feed and thus, require no special care. It is also not a  
57 vector as the adult flies are neither attracted to human habitat nor foods (van Huis et al.  
58 2013). BSF fly larvae being voracious, convert organic waste in a quick fashion into valuable  
59 biomass, restraining bacterial growth and hence, markedly preclude the generation of  
60 offensive odour. Furthermore, the larvae species aerate and dry up the manure which  
61 enhances odour reduction (van Huis et al. 2013). The larvae have been reported to contain  
62 natural antibiotics as well as modify the micro-flora of manure which eventually reduces  
63 harmful bacteria such as *Escherichia coli* 0157:H7 and *Salmonella enterica* (van Huis et al.  
64 2013).

65 Globally, several insect species, including black soldier fly, have been fed in various life  
66 cycle stages to animals (Anankware et al. 2015; Wallace et al. 2017). However, the use of the  
67 black soldier fly larvae as feed for poultry in West Africa is uncommon (Kenis et al. 2014).  
68 This study was therefore undertaken with the aim of ascertaining the impact that black soldier  
69 fly larval meal (BSFLM) would elicit on growth performance, survivability, sensory and  
70 carcass characteristics, haematological and biochemical indices as well as economics of  
71 production of grower-finisher guinea fowl reared under tropical conditions

72

## 73 **Materials and Methods**

### 74 *Study location and conditions*

75 The study was conducted at the guinea fowl resource centre of CSIR-Animal Research  
76 Institute, Katamanso station, Accra where the larval production was also carried out. The  
77 station is in the coastal savannah zone of Ghana with a mean annual rainfall of 730 mm and

78 two rainy seasons namely major (May – mid-July) and minor (mid-August – October).  
79 Generally, there is very little variation in temperature throughout the year. The monthly  
80 temperature ranges between 24.7 (August) and 28 °C (March) with an annual mean of 26.8 °  
81 C. The relative humidity is usually high with values ranging from 65 to 95%. Wind speed  
82 reportedly ranges between 8 and 16 km per hour (Wallace et al. 2012).

### 83 *Production of BSFLM*

84 Two-day old larvae of black soldier flies were inoculated on an unprocessed fruit waste  
85 mixture composed of 60% watermelon, 20% avocado and 20% mango. The larvae were  
86 harvested using a passive sieving system 10 days after inoculation when they were  
87 considered to be at the “white larvae stage” and this was to minimize chitin concentration.  
88 The harvested larvae were kept overnight in a bowl of saw dust in order to allow the  
89 emptying of gut content. They were, then, washed with clean water, dried and milled in a  
90 hammer mill (3000 rpm, 2 mm sieve; KNUST, Kumasi, Ghana). The milled meal was stored  
91 until required for use.

### 92 *Experimental diets, animals and design*

93 The black soldier larvae meal was systematically mixed with other ingredients at specified  
94 concentrations to produce six experimental diets. All the diets were iso-caloric and iso-  
95 nitrogenous and were fed to the keets *ad libitum* including water from eight to eighteenth  
96 weeks of age. The fishmeal component of the experimental diets was replaced with BSFLM  
97 in the following percentage ratio: T1 (Control) – 100% FM: 0% BSFLM, T2 – 80% FM: 20%  
98 BSFLM, T3 – 60% FM: 40% BSFLM, T4 – 40% FM: 60% BSFLM, T5 – 20% FM: 80%  
99 BSFLM, and T6 – 0% FM: 100% BSFLM. The composition and nutrient values for the diets  
100 are shown in Table 1. Two hundred and forty eight-week old grower guinea fowls with mean  
101 live-weight of  $273.2 \pm 10.9$  g were tagged, weighted and randomly assigned to six floor pens.  
102 Each of the concrete floor pens was of 360 x 210 x 420 cm dimension and covered with 5 cm  
103 good quality wood shavings. Each pen was equipped with two bell drinkers, two feeder trays  
104 as well as a florescent bulb for lighting.

### 105 *Chemical analysis*

106 Proximate composition, calcium, phosphorus and gross energy content of the experimental  
107 diets were determined using methods as described in A.O.A.C. (1990). The diets were  
108 analyzed for nitrogen content using the micro-Kjeldahl method (A.O.A.C. 1990).

### 109 *Biochemical and haematological assays*

110 At day 79 (08.00 GMT), which was the last day of the study, before feeding, four guinea  
111 fowls (two males and two females) were randomly selected from each dietary treatment  
112 groupings. Blood was aseptically drawn from the jugular vein with disposable 5 ml plastic  
113 syringe. 2 ml blood was transferred gently into labelled vacutainer tubes containing EDTA  
114 for whole blood count and the remaining 3 ml into the other vacutainer tubes laced with gel

115 and used for blood chemistry assay. Enumeration of erythrocytes and leukocytes were carried  
116 out manually using the procedures described by Samour (2013). Blood was diluted (1:200)  
117 with Natt-Herrick solution, and counting of RBCs was done using Improved Neubauer  
118 haematocytometer. Haemoglobin was determined using spectrophotometer (Cecil 1000  
119 series, England) at 540 nm using Drabkin's solution. Packed cell volume (PCV) was  
120 determined by duplicate capillary tube. The tubes containing blood samples were centrifuged  
121 at 1,200 g in a micro-capillary centrifuge (Model MB) and read with a Hawksley haematocrit  
122 reader. Thin blood smears were stained with Giemsa and examined microscopically under oil  
123 immersion for leukocyte characterization. For each blood smear, a minimum of 200  
124 leukocytes were counted for the determination of differential leukocyte values. For blood  
125 chemistry assay, blood samples were centrifuged at 3,000 rpm for 5 minutes and the sera  
126 used to determine some key lipid, protein and enzyme profiles. These assays were made with  
127 the aid of reagent kits (Spinreact SA, Ctra. Santa Coloma, Spain) and the targeted  
128 biochemical indices quantified using an automatic device, HITACHI 902 (Japan).

### 129 *Sensory evaluation*

130 At the end of the study, six birds (three cocks and three hens) per treatment were slaughtered  
131 and processed to evaluate the impact of the experimental diets on organoleptic properties of  
132 guinea fowl meat. The processing techniques were in accordance with approved methods for  
133 the processing of meat. The breast muscles of the cooked guinea fowl meat were cut into  
134 pieces, cooked with a common recipe and packaged for the assessment. A total of 19 taste  
135 panellists were trained for the organoleptic evaluation. They washed their mouths with water  
136 after tasting each meat sample and assessed attributes which included tenderness, juiciness,  
137 texture, flavour intensity and overall acceptability. Soon after that, they ranked the meat  
138 samples on the Likert scale of 1 – 8 with 1 being the poorest and 8 the best (Teye et al. 2006).

### 139 *Statistical analysis*

140 Data generated were subjected to analysis of variance using Genstat 14th Edition (VSN  
141 International 2011). Data on growth performance of grower-finisher guineas were subjected  
142 to analysis of covariance (ANCOVA) where initial weights of the birds were used as  
143 covariates in the analysis. Means were separated using Duncan's Multiple Range Test. The  
144 results from sensory evaluation were subjected to analysis of variance using SPSS version 17.  
145 The differences were partitioned using the least significant difference (LSD).

146

## 147 **Results**

148 The influence of graded BSFLM replacement of fish meal on growth performance of grower-  
149 finisher guinea fowl is presented in Table 2. The final weight of the birds at the age of 18  
150 weeks significantly differed ( $P < 0.001$ ). Birds that were fed full BSFL (100%) exhibited the  
151 highest live weights and these were markedly higher than the other treatment groups  
152 including the control. The ADG of birds fed these diets were equally found to be significantly

153 (P=0.009) higher compared to the wholly fish meal diet. The inclusion of BSFLM in the diets  
154 seemed to have elicited a positive linear effect on weight gain of the guinea fowls ( $R^2 = 0.91$ )  
155 as shown in Fig. 1. Feed consumption also differed among the treatment groups (P=0.0072)  
156 with diets 3, 4 and 5 exhibiting similar consumption pattern just as those fed the 100% fish  
157 meal (control). The FCR demonstrated similar responses relative to those fed diets 2, 3, 4 and  
158 5 but had significantly (P = 0.0008) higher appreciation when compared to birds fed the  
159 control diet.

160 Black soldier fly larvae meal inclusion did not affect the survivability of the birds as 80 –  
161 100% fishmeal replaced diets demonstrated significantly (P<0.05) higher survivability  
162 compared to the other treatment groups (see Fig. 2). The haematogram assays (Table 3)  
163 showed that full or partial replacement of fish meal with BSF larval meal in guinea fowls  
164 diets did not compromise (P>0.05) the erythropoietic function as well as WBC differentials.  
165 However, increasing BSFLM beyond 20% elicited significantly (P<0.05) higher MCH  
166 concentration. Similarly, graded BSFLM levels did not impact plasma electrolyte, lipid,  
167 metabolites and enzyme concentration in grower-finisher guinea fowls (Table 4).

168 The responses of some carcass characteristics and organs of the birds to the dietary treatments  
169 were similar (P>0.05) except for dressed weight which was significantly (P=0.049) different  
170 (Table 5). Dressed weight was higher (P<0.05) for birds fed diets 40-100% BSFLM but  
171 comparable (P>0.05) to the control diet (100% fishmeal diet). An assessment by both male  
172 and female trained panellists of the impact of the dietary treatments on organoleptic  
173 properties of guinea fowl meat indicated similar (P>0.05) tenderness, juiciness and texture  
174 for all the dietary treatment groups (Table 6). However, meat of birds fed 100% BSFLM-rich  
175 diet was adjudged to have the best flavour generation. Also, including BSF larval meal from  
176 60 to 100% in place of fish meal in growing guinea fowl diets would elicit overall acceptance  
177 just like 100% fish meal inclusion. Lower BSFLM inclusion up to 40% were the least rated in  
178 terms of acceptability. The sensory properties were similarly rated (P>0.05) for meats from  
179 both cocks and hens, except for acceptability which favoured (P<0.05) meat from hens (Fig.  
180 3).

181

## 182 **Discussion**

183 The study was conducted to evaluate the impact of BSFLM on the productive performance  
184 and meat qualities of growing guinea fowls. The experimental diets were formulated to  
185 contain similar energy and protein (Table 1). The feed intake of 58 – 75 g/d/bird observed in  
186 this study was found to be similar to the 63 – 78 g/d/bird reported by Agbolosu and Teye  
187 (2012) but lower than the 130 – 133 g/d/bird reported in by Teye et al. (2000) for growing  
188 guinea fowls. However, the weight gains of 9.2 - 10.5 g/d/bird observed was found to be  
189 slightly higher than the 6.2 – 7.1 g/d/bird Agbolosu and Teye (2012) reported for similar  
190 birds. Differences in diet composition could be responsible for this observation. In this study,  
191 high BSFLM inclusion (60 -100%) in diets, supported growth better than the high fishmeal

192 diets (60-100%). The high weight gain and FCR observed for birds fed (100% BSFLM)  
193 indicate the potential of BSFLM to grow older guinea fowls (8 – 18 weeks) economically.

194 The survival of guinea fowls under intensive system after 8 weeks of age is known to be high  
195 and therefore, the over 85 - 100% survival exhibited in this study is not unusual. Furthermore,  
196 the high survival rate observed for grower-finisher guinea fowls regardless of the protein  
197 source as well as the level of inclusion was similar to the 82.5 – 98.7% reported for similar  
198 birds by Agbolosu and Teye (2012).

199 The determination of hematological as well as biochemical parameters provide valuable  
200 information for the evaluation of the health status of humans and animals though the lack of  
201 reference values for avian blood profile usually restricts its usage (Talebi et al. 2005). The  
202 immune organs such as spleen and thymus gland are important for the maintenance of normal  
203 immune function of animals (Feng et al. 2007; Ravindran et al. 2006; Wallace et al. 2012)  
204 and the lymphoid organ weights are prevalently assessed as a measure of immune status of  
205 poultry (Pope 1991). In this study, the weight of spleen, as well as its index, were similar  
206 ( $P>0.05$ ) for the treatment groups. It can, therefore, be deduced that the inclusion of BSF  
207 larval meal regardless of levels was as good a protein source as fish meal in maintaining the  
208 immune function and status of growing guinea fowls.

209 Table 4 showcased the blood lipids of grower-finisher guinea fowls fed graded BSF larval  
210 meal. High serum cholesterol and triglycerides are reportedly linked to heart disease, stroke  
211 and heart attack (Kaplan and Szabo 1983; Shutler et al. 1987; A.D.A.M. 2005). The results  
212 obtained relative to the blood chemistry profile as well as the heart risk ratios did not present  
213 black soldier fly larval meal as hypercholesterolaemic nor atherogenic agent. Relative to the  
214 control diet which had 100% fish meal as the main protein source, there were no significant  
215 ( $P>0.05$ ) differences in any of the cholesterol profiles determined. Further to this, the  
216 background diet was not high in cholesterol or fat which usually is the case when the  
217 cholesterol potential of a protein source or material is being ascertained (Shutler et al.  
218 1987; Marfo et al. 1990; Wallace et al. 2001; Landi Librand et al. 2007).

219 The results showed that there was no significant ( $P>0.05$ ) variations in the serum  
220 concentration of urea, creatinine nor any of the electrolytes assayed. This is suggestive of the  
221 fact that BSFLM as protein replacement of fish meal would not disrupt the osmolality  
222 likewise the osmotic balance of the blood of the birds neither would it engender disease state  
223 (e.g. diabetes insipidus, hypokalemia, hyperadrenalism, etc.) that would create distortions in  
224 the electrolyte balance with dire consequences (Kaplan and Szabo 1983). Kaplan and Szabo  
225 (1983) have reported that serum creatinine and urea levels yield useful information on the  
226 impairment or dysfunctional state of the kidney. Levels of urea and creatinine are commonly  
227 used markers of renal physiology and pathology and elevated concentrations of these  
228 metabolites indicate nephrotoxicity (Gowda et al. 2010). It can be suggested that the  
229 statistically ( $P>0.05$ ) similar serum creatinine and urea concentrations of BSFLM fed birds  
230 relative to the control diet imply that kidney impairment or dysfunction did not occur.

231 Several biomarkers have been well established and used to investigate the physio-  
232 pathological status of certain vital organs of the body of animals (Abdel-Wareth et al. 2014)

233 and the intact integrity of the organs is markedly amplified by the status of the liver for  
234 instance. The liver is the site of the biosynthesis of most of the plasma proteins of the blood  
235 and thus, the impairment of the hepatic cells would have reflected in the serum proteins  
236 assayed namely total protein, albumin and globulin (Lehninger 1984). In this study, the AST  
237 and ALT concentration which usually become elevated in liver diseased state (Moss et al.  
238 1987) were found to be statistically similar ( $P>0.05$ ) in the BSFLM fed birds just as the  
239 control birds. The non-incidence of any diseased state in the guinea fowls were further  
240 emphasized by the relatively similar response of the ALT/AST (De Ritis) ratios determined  
241 among birds fed the various dietary treatments. The levels of these enzymes coupled by their  
242 relative concentrations in the plasma are always indicative of the incidence of myriad of  
243 diseases. For instance, in toxic or viral hepatitis, ALT is reported to be characteristically as  
244 high as or higher than AST, and the ALT/AST (De Ritis) ratio, which normally is less than 1,  
245 approaches or becomes greater than unity (Moss et al. 1987; Tietz, 1987).

246 Although the final live weights were higher for BSFLM-rich diets compared to fishmeal-rich  
247 diets, dressed weights were found to be similar. This could be attributed to the fewer number  
248 of birds sampled for assessment. The diets however did not show significant effect on the  
249 other organs measured. The results suggest that including BSFLM at all the levels studied  
250 would elicit similar impact as fish meal in terms of texture, juiciness and tenderness of the  
251 carcasses of the birds fed those diets. However, the diet effect on meat flavour and  
252 acceptability was evident with the complete BSFL diets recording the best flavour and  
253 acceptability ratings. Similar to an earlier report (Al-Qazzaz et al. 2016), BSFL inclusion in  
254 diets of laying hens improved appearance, texture, taste and acceptance of eggs. Meat  
255 acceptability is principally influenced by meat flavour and tenderness (Reicks et al. 2012).  
256 Robbins et al. (2003) suggested that the combination of taste and odour, as well as mouth feel  
257 and juiciness, affect flavour perception. Meat flavour and palatability are largely influenced  
258 by the fat content volatiles from lipid sources. Small proportion of oxidized fatty acids from  
259 lipid sources can be sufficient to alter flavour significantly (Belitz et al. 2009). Feed affects  
260 the physico-chemical and organoleptic parameters of meat, including carcass composition,  
261 degree of fattening, fatty acid profile of meat and formation of short branched-chain fatty  
262 acids (Khan et al. 2015) and can therefore be used to improve poultry meat flavour (Fanatico  
263 et al. 2007). Feed supplements including dietary fat source, dl- $\alpha$ -tocopheryl acetate and  
264 ascorbic acid were reported to have influenced the flavour of chicken meat (Jayasena et al.  
265 2013). Birds fed a diet containing 8% herring meal resulted in fishy, unpleasant, rancid, or  
266 stale flavoured raw meat (Poste 1990).

267 In the current study, the meat characteristics measured were not different between hens and  
268 cocks (Fig. 3) except for meat acceptability. The sex of an animal has been reported to  
269 influence the flavour and general acceptability of meat (Crouse et al. 1981). A significant  
270 influence of sex on the fatty acid profile of *longissimus dorsi* (Lorenzo et al. 2013) as well as  
271 a larger infiltration of fat content in females (Horcada et al. 1998) has led to the suggestion  
272 that that the meat of females should be juicier than the meat of males. Forrest (1975) reported  
273 tenderer, juicier, and more flavourful, with higher overall palatability scores of roasted ribs  
274 from steers than roasts from bulls. Vani et al. (2006) explains that variations in nucleotide  
275 content in muscles can be due to the differences in species, breed, age, sex etc. These can

276 result in different levels of flavour precursors, causing variations in the type and  
277 concentration of volatile compounds. In contrast, Franco et al. (2011) reported no significant  
278 difference of meat quality between sexes.

## 279 **Conclusion**

280 The results obtained in this study demonstrated that BSFLM did not cause any physio-  
281 pathological anomalies in the grower guinea fowls used neither would it at any level of  
282 inclusion adversely impact on growth performance. Organ and haematopoietic integrity were  
283 assured regardless of the protein type used in formulating the diets as well as levels of  
284 inclusion. It is observed that BSFLM could replace fishmeal up to 100% in grower-finisher  
285 guinea fowl diet without compromising on the organoleptic attributes of the carcasses.

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## 293 **Conflict of interest**

294 The authors declare that they have no competing interests.

295

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**Table 1, Experimental diets for grower-finisher guinea fowls**

Ingredients	Dietary treatments (FM:BSFLM)					
	100:0	80:20	60:40	40:60	20:80	0:100
Yellow maize	63.4	65.0	65.0	64.0	63.0	63.0
Soybean meal	14.0	11.0	11.0	12.0	13.0	14.0
Fishmeal	3.00	2.40	1.80	1.20	0.60	-
BSFL	-	0.60	1.20	1.80	2.40	3.00
Lysine	0.50	0.50	0.50	0.50	0.50	0.50
Methionine	0.25	0.25	0.25	0.25	0.25	0.25
Wheat bran	15.2	16.0	16.0	16.0	16.0	16.0
Iodated salt	0.2	0.25	0.25	0.3	0.35	0.35
Oyster shells	2.60	2.70	2.70	2.65	2.60	2.60
Dicalcium phosphate	1.00	1.00	1.00	1.00	1.00	1.00
Vit and Min premix	0.30	0.30	0.30	0.30	0.30	0.30
Calculated analyses (%)						
ME (MJ/kg)	15.6	15.7	15.3	15.4	15.4	15.3
Crude protein	11.5	11.4	11.4	11.4	11.3	11.3

\*Vitamin/mineral premix: Vit. A – 800 IU; Vit. D – 500 IU; Vit. E – 2.5 mg; Vit. K – 1 mg; Vit. B2 – 2 mg; Vit. B12 – 0.005 mg; Folic acid – 0.5 mg; Nicotinic acid – 8 mg; Calcium panthotenate – 2 mg; Choline chloride – 50 mg; Manganese – 50 mg; Zinc – 4 mg; Copper – 4.5 mg; Cobalt – 0.1 mg; Iodine – 1 mg; Selenium – 0.1 mg; ME – Metabolizable energy; Vit – vitamin; Min - mineral

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**Table 2, Growth performance of grower-finisher Guinea fowls fed BSFLM**

Parameters	Dietary treatments (FM:BSFLM)						SEM	P-value
	100:0	80:20	60:40	40:60	20:80	0:100		
Initial weight (g)	318	254	228	285	307	292	10.9	0.132
Final weight(g)	960 <sup>b</sup>	897 <sup>c</sup>	880 <sup>c</sup>	974 <sup>b</sup>	1008 <sup>ab</sup>	1029 <sup>a</sup>	16.0	0.000
Ave. Daily gain(g/bird/day)	9.16 <sup>c</sup>	9.19 <sup>c</sup>	9.31 <sup>bc</sup>	9.84 <sup>abc</sup>	10.0 <sup>ab</sup>	10.5 <sup>a</sup>	0.136	0.009
Feed intake	58.0 <sup>a</sup>	69.3 <sup>b</sup>	71.1 <sup>ab</sup>	70.6 <sup>ab</sup>	75.3 <sup>a</sup>	65.1 <sup>bc</sup>	1.46	0.007

(g/bird/day)

FCR 6.34<sup>bc</sup> 7.57<sup>a</sup> 7.64<sup>a</sup> 7.18<sup>ab</sup> 7.52<sup>a</sup> 6.18<sup>c</sup> 0.162 0.0125

<sup>abc</sup>Means in a row with same or no superscripts are not significantly different (P>0.05)

Ave. – Average; FCR – Feed conversion ratio

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**Table 3, Haematogram and leukogram response of grower guinea fowl fed graded BSFLM diets**

Parameter	Dietary treatments (FM:BSFLM)						SEM	P - value
	100:0	80:20	60:40	40:60	20:80	0:100		
RBC(x 10 <sup>6</sup> /L)	3.47	3.15	3.31	3.14	3.24	3.02	0.0660	0.480
PCV (%)	43.8	38.8	46.8	45.3	43.3	43.8	0.825	0.101
MCV (fl)	127	125	142	144	134	146	2.96	0.107
MCH, (pg)	39.0 <sup>b</sup>	39.2 <sup>b</sup>	46.4 <sup>a</sup>	46.0 <sup>a</sup>	42.4 <sup>ab</sup>	47.2 <sup>a</sup>	1.05	0.033
MCHC (%)	29.3	31.4	32.7	31.9	31.7	32.4	0.352	0.0507
WBC (x10 <sup>9</sup> /L)	20.0	15.1	18.0	18.3	17.8	18.2	0.561	0.275
Neutrophils (%)	29.8	23.3	42.0	31.0	34.0	36.3	2.44	0.394
Lym (%)	53.0	62.8	38.3	47.8	52.5	49.8	3.14	0.324
Basophils (%)	1.75	0.250	0.50	1.50	1.00	1.00	0.200	0.260
Eosinophils (%)	12.3	13.5	18.5	18.0	11.3	10.8	1.89	0.658
Monocytes (%)	3.25	0.25	0.75	1.75	1.25	2.25	0.442	0.442
Spleen index	0.50	0.51	0.45	0.45	0.52	0.51	0.0250	0.968

RBC red blood cell, PCV packed cell volume, WBC white blood cell, Lym lymphocytes, MCHC – Mean Cell Haemoglobin Concentration; MCH – Mean Cell Haemoglobin; MCV – Mean Cell Volume

<sup>abc</sup>Means in a row with the same or no superscript are not significantly different (P>0.05)

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**Table 4, Effect of BSFLM on serum concentrations of lipids, electrolytes, metabolites and enzyme**

Parameter	Dietary treatments (FM:BSFLM)						SEM	P - value
	100:0	80:20	60:40	40:60	20:80	0:100		
Total Chol (mmol/L)	3.55	2.78	4.38	2.98	4.18	3.13	0.242	0.331
	2.10	2.14	2.37	1.39	2.06	1.76	0.206	0.854
Triglyceride (mmol/L)								
HDL (mmol/L)	1.22	1.15	1.39	1.95	1.59	1.84	0.116	0.208
LDL (mmol/L)	1.32	0.648	1.90	0.443	2.15	0.855	0.281	0.458
LDL/HDL Ratio	5.61	0.280	2.52	1.42	0.840	1.10	0.611	0.106
Creatinine (mmol/L)	24.8	29.6	32.8	30.5	40.3	38.1	1.88	0.065
Urea (mmol/L)	9.44	8.77	9.48	8.12	8.63	8.52	0.175	0.168
Na <sup>+</sup>	144	99.0	102	117	106	87.4	7.62	0.394
K <sup>+</sup>	3.10	2.50	2.75	2.03	2.83	2.18	0.221	0.779
Cl <sup>-</sup>	122	115	117	119	119	122	1.63	0.864
ALT (μ/L)	6.32	6.14	14.0	8.34	8.49	9.67	1.36	0.600
AST (μ/L)	230	234	270	234	267	232	7.96	0.515
ALP (μ/L)	1360	1857	1651	984	1742	1492	93.1	0.071
GGT (μ/L)	1.50	4.17	2.60	1.30	10.0	0.440	1.51	0.530
D. Bil (μmol/L)	1.33	4.15	4.55	3.42	5.08	5.94	0.901	0.815
T. Bil (μmol/L)	2.66	4.19	6.60	3.39	9.52	3.64	1.21	0.634
Albumin (g/L)	14.5	13.8	15.0	14.6	14.5	14.1	0.330	0.956
Total protein (g/L)	32.9	29.5	40.0	29.0	32.9	27.0	1.76	0.375
Globulin (g/L)	18.4	15.7	25.0	14.4	18.5	12.9	1.78	0.484

*Tot. Chol* total cholesterol, *HDL* high-density lipoprotein, *LDL* low-density lipoprotein, *ALT* alanine transaminase, *AST* aspartate transaminase, *ALP* alanine phosphatase, *GGT* gamma glutamyl transferase, *D. Bil* direct bilirubin, *T. Bil* total bilirubin

<sup>abc</sup>Means in a row with the same or no superscript are not significantly different (P>0.05)

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**Table 5, Effect of BSF larval meal on live weight and some organs of Guinea fowl (g)**

Parameters	Dietary treatments (FM:BSFLM)						SEM	P-Value
	100:0	80:20	60:40	40:60	20:80	0:100		
Live weight	1161	1058	1224	1135	1165	1181	17.8	0.119
Dead weight	1114	1013	1174	1087	1113	1119	17.2	0.150
Blood weight	46.9	44.8	50.4	48.2	52.2	62.1	2.01	0.106
Dressed weight	785 <sup>a</sup>	680 <sup>b</sup>	837 <sup>a</sup>	741 <sup>ab</sup>	773 <sup>ab</sup>	770 <sup>ab</sup>	15.4	0.0490
Dressing	67.7	64.3	68.3	65.3	66.2	65.1	0.524	0.161
Head	36.4	36.0	39.8	39.6	40.1	39.3	0.806	0.578
GIT <sup>†</sup>	42.0	43.7	47.1	45.3	42.5	44.3	1.27	0.862
Heart	6.70	6.05	6.33	7.05	7.98	7.05	0.282	0.217
Liver	11.7	13.1	12.2	12.5	13.1	15.5	0.578	0.550
Gizzard	23.4	32.4	24.8	25.9	25.9	22.3	1.26	0.0825
Spleen	0.575	0.525	0.550	0.500	0.600	0.600	0.0260	0.885

<sup>ab</sup>Means in a row with the same or no superscript are not significantly different (P>0.05)

<sup>†</sup>GIT – Gastro intestinal tract

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**Table 6, Impact of dietary treatments organoleptic properties of guinea fowl meat (Likert scale)**

Organoleptic properties	Dietary treatments (FM:BSFLM)						SEM	P-value
	100:0	80:20	60:40	40:60	20:80	0:100		
Tenderness	4.92	5.21	5.08	4.97	4.58	5.37	0.104	0.345
Juiciness	4.50	4.87	4.89	4.95	4.79	5.06	0.0776	0.386
Texture	4.53	4.87	5.03	5.05	4.50	4.95	0.0856	0.228
Flavour	4.39 <sup>a</sup>	4.39 <sup>a</sup>	5.05 <sup>b</sup>	4.29 <sup>a</sup>	4.92 <sup>a</sup>	5.18 <sup>b</sup>	0.0968	0.017

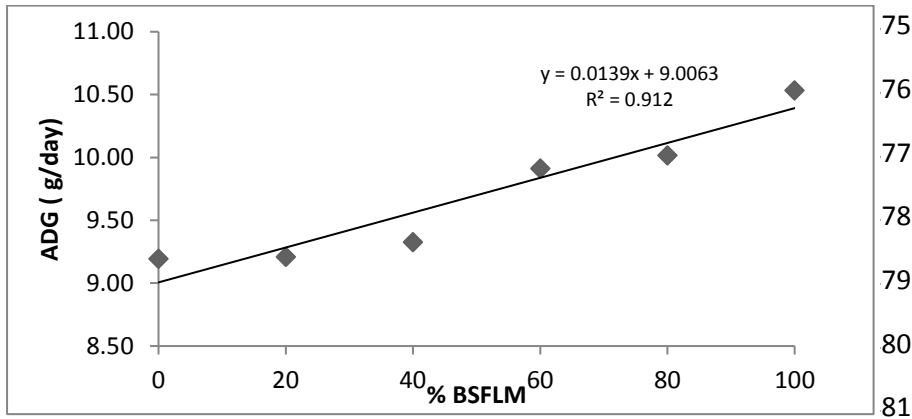
Acceptability 4.75<sup>ab</sup> 2.75<sup>a</sup> 3.00<sup>a</sup> 5.14<sup>b</sup> 5.32<sup>b</sup> 5.66<sup>b</sup> 0.148 0.001

<sup>ab</sup>Means in a row with the same or no superscript are not significantly different (P>0.05)

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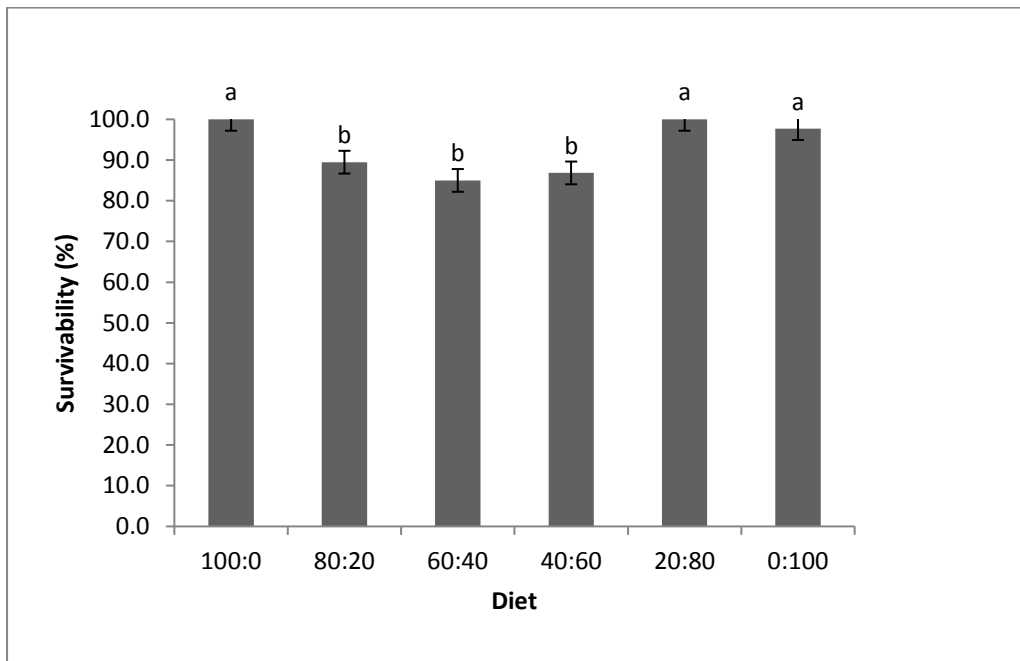
**Fig. 1. Regressional analysis of fish meal replacement with BSFLM relative to ADG** [P = 0.0030; S.E. = 0.180]

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**Fig. 2. Survivability of grower-finisher guinea fowls fed diets containing varying levels of BSFLM**

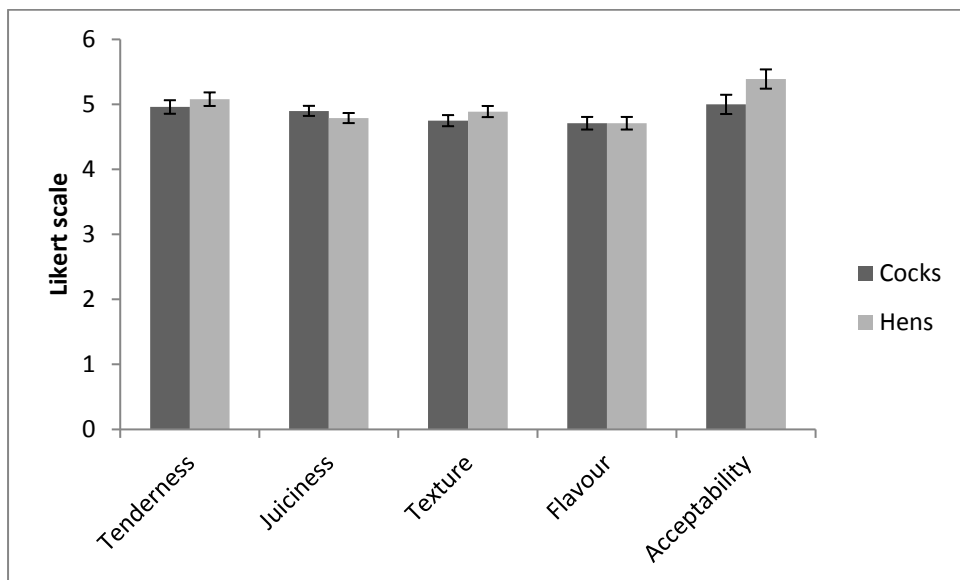


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**Fig. 3. Sensory evaluation of guinea fowl meat from cocks and hens**

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