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#### 3 Abstract

4 This study investigates the bioconversion efficiency and nutrient accumulation in black soldier fly (BSF; 5 Hermetia illucens), focusing on the effects of feeding on two different substrates (tofu by-products and 6 food waste) and harvesting at two developmental stages (larvae and prepupae) within a 2×2 factorial 7 arrangement. The growth performance, conversion efficiency, nutrient composition, amino acid profile, 8 fatty acid composition, and nutrient composition of BSF meal were assessed. Results indicated that BSF 9 reared on tofu by-products exhibited superior weight gain compared to those fed food waste, with significant enhancements observed in weight, length, and width upon harvesting at the prepupae stage. 10 Moreover, tofu by-products promoted higher bioconversion rates, protein conversion efficiency, and 11 lipid yield, while food waste favored lipid conversion. Analysis of nutrient composition revealed higher 12 crude protein content in BSFs fed tofu by-products, with elevated levels of crude protein, ether extract, 13 and chitin in prepupae-stage BSFs. Higher concentrations of isoleucine, leucine, and tryptophan were 14 observed in tofu by-product-fed BSF. Conversely, BSFs harvested at the prepupae stage exhibited 15 increased levels of threonine, alanine, and tyrosine, regardless of substrate. Higher proportions of  $\alpha$ -16 linolenic acid (C18:3n3) and docosahexaenoic acid (C22:6n3) were observed in tofu by-product-fed 17 BSF. Conversely, BSF harvested at the larval stage displayed higher levels of saturated fatty acids, 18 including lauric acid (C12:0) and myristic acid (C14:0). In conclusion, tofu by-products emerged as a 19 20 promising substrate for enhancing essential amino acid and unsaturated fatty acid content in BSF, while 21 harvesting at the prepupae stage offered advantages in nutrient density and storage stability of the 22 harvested biomass.

Keywords: Food waste, tofu, bioconversion, larvae, prepupae, black soldier fly 23

#### **25 INTRODUCTION**

26 Sustainable agriculture strives to meet the world nutritional needs while minimizing environmental 27 degradation and resource depletion [1,2]. Insects have emerged as promising candidates owing to their environmental sustainability and efficiency [3]. Among these alternatives, the black soldier fly (BSF; 28 Hermetia illucens) has garnered considerable attention in recent years due to its potential to address 29 sustainability challenges in agriculture [4,5]. The larvae of BSF offer numerous advantages as a protein 30 31 source for animal nutrition, offering a viable solution to food security concerns. A primary advantage lies in BSF remarkable efficiency in converting organic waste into protein biomass [4,6,7]. These 32 33 voracious larvae possess the ability to decompose a wide array of organic substrates such as food waste, agricultural residue, and animal manure into nutrient-rich biomass [2,3,8], presenting a sustainable 34 solution to waste management while augmenting feed production. The larvae emit fewer greenhouse 35 gases, rendering BSF cultivation a more eco-friendly alternative to traditional livestock farming 36 37 practices [9].

Economically, BSF production offers cost-effective advantages over conventional livestock rearing [7].
Notably, by-products from tofu processing, such as tofu whey, represent an underutilized feed resource
for BSF rearing [3]. The abundance of tofu waste, estimated at millions of tons annually in select regions,
underscores the potential for BSF cultivation to mitigate waste disposal challenges while generating
valuable protein biomass [3].

Nevertheless, the nutritional composition and growth performance of BSF larvae are influenced by their diet, particularly botanical waste substrates [10–12]. Optimization of dietary inputs is pivotal to maximizing protein production efficiency per unit of organic waste, thereby enhancing the sustainability and cost-effectiveness of BSF-based feed production [8]. Moreover, the nutrient composition of BSF meal can be under the influence of the growth stage harvest including larvae and prepupae [13]. Thus, a systematic evaluation of BSF's growth dynamics, conversion efficiency, and larval quality across different botanical waste substrates is imperative to identify optimal feeding regimes. This study endeavors to assess the growth performance, conversion efficiency, and larval quality of BSF reared on various botanical waste substrates, namely food waste and tofu by-products. By elucidating the effects of substrate type and developmental stage (larvae or prepupae) on nutrient composition, weight, length, development time, substrate consumption index, conversion efficiency, and fatty acid composition, this research aims to contribute to the optimization of BSF rearing practices for sustainable and efficient protein production in the context of waste valorization and animal feed supplementation.

57

#### 58 MATERIAL AND METHODS

#### 59 Larvae rearing control

The experiment was conducted at a commercial BSF farm located in Eui-sung, Gyeongsangbuk-do, 60 61 Republic of Korea. Prior to hatching, the BSF insects used in this study were sourced from the farm. 62 The trial colony was meticulously maintained under controlled environmental conditions, with a temperature set at 27±0.3°C, relative humidity maintained at 65±8%, and a photophase of 16 hours of 63 natural light supplemented with LED lighting providing an intensity of 6000 lux. For the experimental 64 setup, a total of 30 egg clutches were utilized, each containing approximately 50,000 larvae. The 65 66 average weight of the egg clutches was measured at 24.16±0.91 mg per clutch, with an average egg weight of 0.027 mg. The egg clutches were further divided into four treatment groups, comprising two 67 different substrates (tofu by-product and food waste; Table 1) and two distinct growth stages (larvae 68 and prepupae). Each treatment group consisted of 10 replicates, housed within experimental boxes 69 70 measuring 1,064×507×320 mm (length×width×height). The trial continued until the end of the prepupal 71 stage, just prior to the pupal stage transition. Egg counting within each clutch was performed according 72 to the methodology outlined by Georgescu et al. c, employing an Alpha model binocular magnifier with 73 7x to 45x magnification. To ensure accuracy, the eggs were dispersed in 70% ethanol and subsequently 74 captured and enumerated through photographic means utilizing the ClickMaster2000 1.0 software,

- 75 accessible at https://www.thregr.org/~wavexx/. This meticulous process facilitated precise egg counting,
- ressential for subsequent experimental procedures and data analysis.

## 77 Assessment of larval growth performance

To assess the growth performance of BSF larvae, measurements were taken at the time of harvest, including weight, length, width, and survivability. In order to standardize the comparison across harvest days, 100 larvae were randomly selected for measurement beginning on day 10, and continuing until the average larval weight reached 190 mg. The larval yield, indicative of survivability on the harvest day, was calculated using the following equation [3]:

Larval yield % = 
$$\frac{\text{Larvae}_F}{\text{Larvae}_I} \times 100$$

83 Where:

Larvae<sub>F</sub> represents the number of larvae at the end of the trial. Larvae<sub>I</sub> represents the number of larvae
at the initiation of the trial.

This approach allowed for a comprehensive evaluation of larval growth and survival dynamics, essential for assessing the efficacy of different experimental treatments. Additionally, the utilization of standardized measurement protocols ensured accuracy and reliability in the assessment of larval performance across varying experimental conditions.

## 90 Substrates preparation

The substrates utilized in this study were procured from local farms and facilities to ensure relevance to regional waste streams. Prior to their use, dried substrates were moistened with water to achieve a target moisture content of 65%. Egg blocks were then positioned on two distinct substrates: food waste and tofu by-products. The food waste substrate consisted of organic waste collected from households, kitchens, and local dining establishments through a municipal waste collection facility located in Euisung, Gyeong-sangbuk-do, Republic of Korea. Tofu by-products, comprising soybean curd residue, were sourced from a local tofu manufacturing facility situated in Kang-reung, Kangwon-do, Republic

98 of Korea. To prevent fermentation and ensure uniformity across substrates, the materials were promptly dried using an industrial drying apparatus (Model: KAPD-A098D, Manufactured in Gwangju, Republic 99 100 of Korea), operating at a temperature of 60°C for 48 hours, until the moisture content was reduced to less than 8%. Upon commencement of the experiment, water was applied to the substrates to reach the 101 102 desired moisture level of 65%. Each experimental box was provisioned with a total of 20 kg (dry matter 103 basis) of substrate. To create a suitable environment for larval development and prevent drowning, a 104 layer of 1 kg of recycled paper pulp and 600 g of sawdust was spread atop the substrates. This bedding 105 layer facilitated larval movement and provided an additional substrate for microbial activity, which may 106 contribute to the decomposition process.

#### 107 Chemical composition analysis

The chemical composition of the harvested larvae from each treatment was determined to assess their nutritional 108 109 profile. The contents of dry matter (DM), crude protein (CP), ether extract (EE), crude ash (Ash), and chitin were 110 analyzed using standardized methods outlined by the Association of Official Analytical Chemists [14]. Initially, fresh larvae were weighed to establish baseline measurements. Subsequently, the larvae were subjected to drying 111 112 in an oven set at 85°C for a duration of three days to determine the DM content, following the procedure outlined 113 in method 930.15 of the AOAC [14]. To quantify the protein content, AOAC official method 990.03 was 114 employed, utilizing the Kjeldahl method. This method entails the digestion of the sample with sulfuric acid to 115 release nitrogen, followed by conversion of the nitrogen to ammonia. The ammonia is then distilled and titrated 116 to determine the nitrogen content, which is subsequently utilized to calculate the protein content [14]. Similarly, 117 the ether extract content was determined using AOAC official method 990.15, involving the extraction of lipid 118 content from the sample with a suitable solvent, followed by evaporation of the solvent to obtain the lipid content 119 [14]. The gross energy of BSF larvae was measured using a bomb calorimeter (Model 1261, Parr Instrument Co., 120 Moline, IL, USA), providing insight into the calorific value of the larvae. Following analysis, any remaining 121 samples were promptly stored at  $-20^{\circ}$ C for future analysis and reference, ensuring preservation of sample integrity 122 for subsequent investigations. The amino acid (AA) composition of BSF meal was assessed using highperformance liquid chromatography (HPLC) with a Waters 486 system (Waters, Milford, MA, USA). 123 124 This analysis was conducted following acid hydrolysis, as described by Lee et al. [15], to release the 125 amino acids from the protein matrix for quantification. To ensure comprehensive analysis, the determination of methionine and cysteine was performed after oxidation with performic acid, followingthe methodology outlined by Kim et al. [16].

## 128 Larval convergence efficiency evaluation

To assess the convergence efficiency of larvae reared on different botanical wastes, several parameters including bioconversion rate (BCR), waste reduction rate (WR), protein conversion rate (PCR), lipid conversion rate (LCR), protein yield (PY), lipid yield (LY), and mass distribution patterns were evaluated using the following equations.

133 The BCR was calculated as:

134 BCR % = 
$$\frac{DM_F - DM_I}{DM_S} \times 100$$

135

 $\label{eq:matter} 136 \qquad \mbox{Where } DM_F \mbox{ represents the dry matter of BSF larvae at harvest day, } DM_I \mbox{ represents the dry matter of } \\$ 

137 BSF larvae at the initiation of the trial, and DMS represents the total dry matter of the substrate.

138 The WR was determined by:



- 139 Where  $DM_S$  represents the total dry matter of the substrate and  $DM_R$  represents the residual dry matter 140 of the substrate after larval consumption.
- 141 The PCR was calculated as:

$$PCR \% = \frac{DM_F \times Protein_L \%}{DM_S} \times 100$$

- 142 Where protein<sub>L</sub> % represents the protein content in the larvae, DM<sub>F</sub> represents the dry matter of BSF
- 143 larvae at harvest day, and DMS represents the total dry matter of the substrate.
- 144 The LCR was determined by:

$$LCR \% = \underbrace{\frac{DM_F \times EE_L \%}{DM_S}}_{DM_S} \times 100$$

145 Where  $EE_L$  % represents the lipid content in the larvae,  $DM_F$  represents the dry matter of BSF larvae at 146 harvest day, and  $DM_S$  represents the total dry matter of the substrate.

147 The PY was calculated as:

$$PY kg/c = g of DM_{F yield} \times CP_L \%$$

148 Where DM<sub>F</sub> yield represents the dry matter of BSF larvae at harvest, and CP<sub>L</sub> % represents the crude

149 protein content in the larvae.

150 The LY was determined as:

$$LY \text{ kg/c} = g \text{ of } DM_{F \text{ yield}} \times EE_L$$

151 Where  $DM_F$  yield represents the dry matter of BSF larvae at harvest,  $EE_L$  % represents the ether extract 152 content in the larvae, and 'c' represents the clutch size (50,000 larvae).

153

#### 154 Fatty acid composition

155 The determination of fatty acid content was conducted following the methodology outlined by Georgescu et al. [17]. For analysis, 10 g of meal sample was collected and subsequently placed on a 156 clean substrate for a 24-hour period. Following this, the samples were washed with 70% ethanol to 157 remove any impurities and then stored at -80°C to maintain sample integrity. Extraction and 158 159 identification of fatty acid methyl esters (FAME) from BSF larvae fats were performed using gas chromatography with mass spectrometry detection, adhering to established standards including AOAC-160 969.33 [14], ISO 3657: 2002, ISO 12966-2: 2011, and ISO 12966-2: 2017. The analytical process 161 162 involved saponification of the fat, achieved by subjecting the samples to methanolic sodium hydroxide 163 solution (0.5 M) at 210°C on a sand bath with a reflux rate of 1 drop/s. Subsequently, esterification with

164 a 15% vol boron trifluoride catalyst and cooling with hexane were carried out. Gas chromatographic analysis was conducted using a Perkin Elmer chromatographic system equipped with a mass 165 166 spectrometer detector (GC-MS). This system featured an Elite-Wax chromatographic column with a stationary polar phase polyethylene glycol measuring 30 m in length, 0.25 mm in internal diameter, and 167 a film thickness of 1.0 μm. Operating parameters included an injection port temperature of 220°C, an 168 injected sample volume of 1.0  $\mu$ L, a helium carrier gas flow rate of 1.5 mL/min, and a splitting ratio of 169 170 40:1. The mass spectrometer settings comprised a transfer line temperature of 150°C, a source temperature of 150°C, a multiplier of 1500, and a solvent delay of 0-1.5 min. Quantification of fatty 171 acid concentration was achieved by comparing the relative retention time of FAME to the certified 172 standard Mix FAME Supelco. The individual fatty acid concentrations were expressed as a percentage 173 of the total identified FAME. Moreover, key ratios between various fatty acid categories, including total 174 saturated fatty acids (SFA) and total unsaturated fatty acids, were calculated. Additionally, the ratios 175 between fatty acids from the n-6 and n-3 series were determined to further elucidate the nutritional 176 profile of the BSF biomass. 177

#### 178 Statistical analysis

179 The data were subjected to analysis employing a 2×2 factorial arrangement utilizing the General Linear Model procedure of SAS (version 9.4, 1996, SAS Inst. Inc., Cary, NC, USA). Differences in various 180 parameters including growing performance, conversion efficiency, nutrient composition, amino acid, 181 and fatty acid concentrations among the treatments were assessed using a two-way ANOVA test. To 182 183 further elucidate the differences between treatments, the Tukey test was employed to compare the means of the treatments. Statistical significance was determined at a threshold of p < 0.05. Moreover, a p-value 184 less than 0.1 was considered indicative of a trend. The results are expressed as the mean  $\pm$  standard 185 186 deviation, providing insights into the central tendency and variability of the data.

187

188 RESULTS

#### 189 Growth performance

The BSF larvae reared on tofu by-products exhibited significantly higher (p < 0.05) weight gain 190 191 compared to those fed with food waste (Table 2). Moreover, harvesting at the prepupae stage resulted in notable enhancements in weight, length, and width of BSF, indicating superior growth performance 192 193 compared to larvae-harvested BSF, which displayed higher survival rates. The BSF larvae reared on tofu by-products exhibited significantly higher (p < 0.01) BCR, PCR, PY, and LY compared to those 194 195 fed with food waste (Table 2), however, LCR was lower in BSF fed tofu by-products compared with 196 food waste. Moreover, harvesting at the prepupae stage resulted in lower BCR compared to larvae harvested BSF, however, harvesting at the prepupae stage showed higher PY, LCR, PY, and LY 197 198 compared to larvae harvested BSF.

199

#### 200 Nutrient composition of meal

The analysis revealed no significant differences in dry matter, gross energy, or ash content across different substrates and developmental stages (Table 3). However, BSFs fed on tofu by-products exhibited a higher crude protein content compared to those fed with food waste. Furthermore, harvesting at the prepupae stage led to elevated levels of crude protein, ether extract, and chitin, indicating enhanced nutrient composition in prepupae-stage BSF.

#### 206 Amino acid profile of the meal

Amino acid profiling demonstrated distinct compositions in BSF fed with different substrates and harvested at different developmental stages (Table 4). BSFs reared on tofu by-products showed increased concentrations of isoleucine, leucine, tryptophan, alanine, and proline. In contrast, BSFs harvested at the prepupae stage displayed higher levels of threonine, tryptophan, alanine, serine, and tyrosine, indicating variations in amino acid profiles influenced by both substrate type and developmental stage.

#### 213 Fatty acid composition of meal

Fatty acid analysis revealed variations in the composition of BSF lipids based on substrate and developmental stage (Table 5). BSFs fed on tofu by-products exhibited higher ratios of C10:0, C12:0, C14:0, C14:1, C18:3n3, and C20:1 fatty acids, while those consuming food waste showed a higher content of C18:4. Additionally, SFA such as C12:0, C15:0, C17:0, and C18:1 were more abundant in BSFs harvested at the prepupae stage.

## 219 DISCUSSION

220 The higher weight gain observed in BSF larvae reared on tofu by-products compared to those fed with 221 food waste suggests that the nutrient composition and bioavailability in tofu by-products are more conducive to larval growth. The higher protein in tofu by-products may facilitate improved growth rates. 222 The enhancements in weight, length, and width of BSF when harvested at the prepupae stage could be 223 attributed to the larval maturation process. As larvae transition into prepupae, they are in a state of 224 225 maximal nutrient accumulation and tissue development [7]. This stage is also characterized by efficient energy storage and nutrient conversion in preparation for metamorphosis [13], which may explain the 226 observed improvements in growth performance at this developmental stage. In contrast, the higher 227 survival rates in larvae harvested BSF as opposed to those harvested at the prepupae stage may be linked 228 to the life cycle and physiological state of the larvae. Larval stages possess more resilience and 229 adaptability to varying environmental conditions and nutrient availability compared to prepupae [18]. 230 This higher adaptability results in increased survival rates when harvesting at the larval stage [8]. 231 232 Overall, these results highlight the importance of substrate choice and timing of harvest in optimizing 233 BSF growth performance. The mode of action related to nutrient availability, larval development, and life cycle dynamics are central to understanding the differences observed in growth performance. 234

The lack of significant differences in dry matter, gross energy, and ash content across different substrates and developmental stages suggests that the overall energy density and mineral content of BSF biomass remain relatively consistent regardless of dietary inputs or harvest timing. However, BSF fed tofu byproducts demonstrated higher crude protein content compared to those fed food waste. This difference can be attributed to the higher protein levels in tofu by-products [3]. Harvesting BSF at the prepupae stage led to increased levels of crude protein, ether extract, and chitin. This suggests that as BSF larvae
mature and transition to the prepupae stage, they undergo biochemical and physiological changes that
enhance their nutrient composition [19]. The increase in crude protein and ether extract may be due to
the prepupae preparation for metamorphosis, which involves accumulating energy and nutrient reserves.
Additionally, elevated chitin levels are associated with the development of the exoskeleton as the insects
approach the pupal stage [20].

246 The amino acid profile of BSF is a crucial aspect of its nutritional value and potential applications as animal feed [3,13]. BSF reared on tofu by-products exhibited increased concentrations of isoleucine, 247 leucine, tryptophan, alanine, and proline compared to those fed with food waste. This suggests that tofu 248 by-products provide a rich source of essential and conditionally essential amino acids, contributing to 249 250 a more balanced and nutritious profile for the larvae. The higher levels of these amino acids may result from the nutrient-rich nature of tofu by-products, which could provide an optimal blend of proteins and 251 other macronutrients for larval development. In contrast, BSF harvested at the prepupae stage displayed 252 elevated levels of threonine, tryptophan, alanine, serine, and tyrosine compared to those harvested at 253 the larval stage. This shift in amino acid composition during the prepupae stage is linked to the 254 physiological changes the insects undergo in preparation for metamorphosis [7,13,21]. The variation in 255 amino acid profiles based on substrate type and developmental stage underscores the importance of 256 optimizing these factors for specific applications of BSF biomass. 257

258 The fatty acid composition of BSF is a critical aspect of its nutritional quality and potential applications 259 in various industries, including animal feed and biodiesel production [2,13]. BSF fed on tofu by-260 products exhibited higher ratios of specific fatty acids, including C10:0, C12:0, C14:0, C14:1, C18:3n3, and C20:1, compared to those consuming food waste. These differences in fatty acid composition may 261 be due to the lipid profiles present in the respective substrates. The observed higher content of C18:4 in 262 263 BSF consuming food waste suggests preferential utilization or accumulation of this fatty acid in 264 response to the substrate lipid content. Additionally, SFA such as C12:0, C15:0, C17:0, and C18:1 were more abundant in BSF harvested at the prepupae stage. This is attributed to metabolic changes 265

associated with larval development and maturation into prepupae [19]. As the larvae transition to the prepupal stage, they undergo metabolic shifts to support pupation and subsequent adult emergence [2,19], leading to alterations in fatty acid synthesis and accumulation. The variations in the fatty acid composition of BSF biomass have significant implications for its nutritional value and suitability for various applications [22]. For instance, certain fatty acids, such as omega-3 and omega-6 fatty acids, are essential for animal health [1,2,11] and contribute to the nutritional quality of BSF-based feed formulations.

BSF meal derived from larvae reared on tofu by-products exhibited higher gross energy compared to 273 meal from food waste-fed BSF. This difference could be attributed to the higher energy density of the 274 tofu by-product substrate, which may have contributed to enhanced energy accumulation in the larvae. 275 276 As a result, the meal from tofu by-product-fed BSF could provide a more energy-rich feed ingredient. Moreover, meals from prepupae-stage BSF demonstrated enhanced gross energy, crude protein, and 277 chitin content. These changes can be due to the physiological and biochemical transformations that 278 occur as the larvae mature into prepupae [2,6,19]. The increased gross energy may result from the 279 280 accumulation of energy reserves to support metamorphosis, while the higher crude protein and chitin 281 levels are associated with tissue development and exoskeleton formation during the pupal transition [2,20]. The distinct fatty acid profile favoring C12:0, C18:3n3, and C22:6n3 in meals from prepupae-282 283 stage BSFs suggests alterations in fatty acid metabolism as the larvae approach the pupal stage. These 284 specific fatty acids may be selectively synthesized or accumulated in response to developmental cues, contributing to the observed fatty acid composition. The higher levels of SFA in prepupae-stage BSF 285 meal could reflect shifts in metabolic pathways aimed at supporting pupation and adult emergence 286 [9,10]. Saturated fatty acids play a role in providing stable energy sources during this critical 287 288 developmental phase [2,10,17]. Understanding the mode of action underlying these variations in 289 nutrient composition can help optimize BSF meal production for targeted applications. The higher gross 290 energy and crude protein content in prepupae-stage BSF meal increases its value as a feed ingredient 291 for high-performance livestock.

### 292 Conclusion

Our results demonstrate that substrate type significantly influences the growth performance, nutrient 293 composition, and fatty acid profile of BSF larvae and prepupae. Larvae reared on tofu by-products 294 exhibited superior growth performance and higher levels of crude protein, while those fed food waste 295 displayed distinct fatty acid compositions. Additionally, harvesting BSF at the prepupae stage resulted 296 297 in enhanced nutrient composition and fatty acid profiles, indicating the importance of the developmental 298 stage in optimizing biomass quality. Furthermore, it was revealed that by understanding the mode of 299 action underlying these variations, we can develop strategies to optimize BSF production systems for 300 farm animal nutrition, sustainable agriculture, and waste management.

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## 374 TABLES

Substrate (%)Food wasteTofu By-productDry matter16.9523.96Crude protein19.6830.52Ether extract21.6512.95Ash6.857.96

Table 1. Nutrient composition of botanical waste used in trial.

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Table 2. Effects of substrate (SUB) and development stage (STG) on growth performance of black soldier fly.

Treatmonte	Food waste		Tofu by-product		SEM		P-value		
	Larvae	Prepupae	Larvae	Prepupae	SEIVI	SUB	STG	SUB×STG	
Larvae growth									
Weight, mg	177.92	191.05	194.97	209.39	2.178	< 0.001	< 0.001	0.802	
Length, mm	13.83	15.20	14.20	15.21	0.132	0.305	< 0.001	0.342	
Width, mm	2.79	2.94	2.84	2.95	0.021	0.477	0.002	0.614	
Survival rate, %	80.14	77.53	77.97	75.50	0.575	0.056	0.022	0.948	
Substrate consumption, %	69.01	77.65	72.16	79.55	0.971	0.090	< 0.001	0.671	
Conversion efficiency rate, %									
Bioconversion	12.10	11.61	12.67	12.36	0.084	< 0.001	0.001	0.440	
Protein conversion	6.12	6.93	9.91	11.02	0.040	< 0.001	< 0.001	0.069	
Lipid conversion	5.88	6.64	4.45	4.96	0.025	< 0.001	< 0.001	0.082	
Protein yield, kg/c	1.13	1.32	1.32	1.47	0.029	0.008	0.005	0.859	
Lipid yield, kg/c	1.03	1.16	1.13	1.26	0.022	0.039	0.006	0.902	

377 SEM, standard error of mean. Each value is the average of 10 replicates.

Treatments	Food waste		Tofu by-product		SEM	P-value		
	Larvae	Prepupae	Larvae	Prepupae		SUB	STG	SUB×STG
DM, %	40.60	42.73	41.57	42.97	0.689	0.662	0.207	0.792
GE, kcal	4,315	4,343	4,388	4,410	30.0	0.253	0.684	0.964
CP, % of DM	39.12	41.66	41.63	43.41	0.326	0.002	0.002	0.569
EE, % of DM	35.53	36.67	35.76	37.02	0.283	0.617	0.041	0.920
Ash, % of DM	7.04	7.24	7.10	7.29	0.167	0.872	0.564	0.916
Chitin, % of DM	7.85	8.45	8.15	8.60	0.125	0.257	0.049	0.882

Table 3. Effects of substrate (SUB) and development stage (STG) on nutrient composition of black soldier fly.

SEM, standard error of mean; DM, dry matter; GE, gross energy; EE, ether extract. Each value is the 378

379 average of 10 replicates.

Treatments	Food waste		Tofu by-product		SEM	P-value					
	Larvae	Prepupae	Larvae	Prepupae		SUB	STG	SUB×STG			
Essential amino acid, %											
Arginine	1.86	1.95	1.74	1.93	0.036	0.361	0.061	0.520			
Histidine	1.06	1.07	1.19	1.11	0.028	0.296	0.062	0.710			
Isoleucine	1.45	1.16	1.66	1.83	0.031	0.002	0.016	0.958			
Leucine	2.39	2.44	2.53	2.68	0.030	0.005	0.114	0.426			
Lysine	2.81	2.92	2.84	2.93	0.062	0.836	0.423	0.931			
Methionine	0.59	0.64	0.54	0.58	0.020	0.219	0.321	0.951			
Phenylalanine	1.07	1.21	1.14	1.23	0.033	0.109	0.164	0.459			
Threonine	1.36	1.48	1.38	1.58	0.022	0.205	0.002	0.400			
Tryptophan	1.97	2.21	2.23	2.32	0.030	0.007	0.015	0.243			
Valine	2.53	2.58	2.59	2.74	0.076	0.479	0.513	0.716			
Non-essential a	mino acid	l, %									
Alanine	2.46	2.56	2.63	2.77	0.029	0.004	0.049	0.680			
Aspartate	3.16	3.49	3.34	3.46	0.056	0.522	0.054	0.355			
Cysteine	0.26	0.28	0.31	0.32	0.018	0.177	0.747	0.927			
Glutamate	3.92	4.02	3.82	4.01	0.175	0.863	0.684	0.893			
Glycine	1.93	2.06	2.11	2.13	0.031	0.062	0.243	0.335			
Proline	2.50	2.64	2.72	2.73	0.026	0.006	0.183	0.203			
Serine	1.37	1.58	1.42	1.55	0.025	0.894	0.002	0.449			
Tyrosine	1.97	2.23	2.14	2.25	0.024	0.070	0.001	0.117			

Table 4. Effects of substrate (SUB) and development stage (STG) on amino acid profile of black soldier fly.

381 SEM, standard error of mean. Each value is the average of 10 replicates.

Treatments	Food waste		Tofu by-product		SEM	P-value		
Iteatilients	Larvae	Prepupae	Larvae	Prepupae		SUB	STG	SUB×STG
Fatty acid, %								
C10:0	0.88	0.91	0.94	0.97	0.009	0.006	0.160	0.827
C12:0	22.22	23.28	23.72	24.90	0.174	< 0.001	0.004	0.871
C14:0	4.38	4.63	4.71	4.68	0.042	0.035	0.238	0.111
C14:1	0.12	0.10	0.16	0.15	0.010	0.032	0.437	0.814
C15:0	0.14	0.16	0.16	0.20	0.008	0.051	0.033	0.454
C16:0	15.43	16.36	14.64	14.35	0.335	0.050	0.641	0.373
C16:1	3.25	3.31	3.29	3.25	0.017	0.882	0.730	0.125
C:17:0	0.22	0.26	0.19	0.23	0.007	0.065	0.015	0.861
C18:0	1.90	2.27	2.03	2.14	0.045	0.985	0.016	0.160
C18:1	24.10	23.41	23.89	23.06	0.268	0.607	0.174	0.893
C18:2n6	19.25	17.04	17.86	17.51	0.328	0.496	0.065	0.170
C18:3n3	2.16	2.11	2.23	2.25	0.022	0.036	0.811	0.392
C18:4	0.90	0.91	0.70	0.74	0.007	< 0.001	0.105	0.527
C20:1	0.24	0.27	0.45	0.47	0.010	< 0.001	0.345	0.811
C20:4n6	0.26	0.21	0.22	0.21	0.009	0.290	0.067	0.252
C20:5n3	1.36	1.28	1.23	1.31	0.017	0.121	0.980	0.028
Other	2.01	2.14	2.35	2.30	0.049	0.016	0.685	0.374
SFA	45.17	47.86	46.39	47.46	0.400	0.612	0.030	0.325
USFA	53.64	50.77	52.38	51.25	0.417	0.647	0.026	0.311

Table 5. Effects of substrate (SUB) and development stage (STG) on fatty acid composition of black soldier fly.

383 SFA, saturated fatty acids; USFA, unsaturated fatty acid. Each value is the average of 10 replicates.