1	Original Research Article
2 3	Effects of using different types of organic wastes for the mass culture of Moina
4	macrocopa
5 6	ABSTRACT

7 Moina macrocopa was cultured with different animal manures (chicken manure, Pig manure and cow 8 manure) and food waste to determine the impact of these food sources on its mass production. All 9 diets were provided at five different concentrations: 500, 1000, 2500, 5000 and 10000 ppm. Gross and 10 net reproductive rates were higher in 1000 ppm concentration of food waste medium and the highest 11 average population growth was obtained of about 9 individual M. macrocopa per ml, whereas pig 12 manure treatment showed the lowest among all the culture medium. Highest population density was 13 observed in low concentration treatments, on the contrary higher concentrations showed adverse effect on *M. macrocopa* cultivation. The results of this study suggest that 1000 ppm concentration of 14 15 food waste produces better results than other animal manures which showed the highest population 16 density and exhibited comparatively higher percentage of highly unsaturated fatty acids than the other 17 treatments and could be an inexpensive and sustainable cultivation approach of Moina macrocopa. 18 Keywords: Moina macrocopa, Animal manure, Food waste, Vial test, Life table demography

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To feed the increasing human population, it is therefore imperious to upgrade aquaculture, including fish farming whose development goes essentially through the success of larval rearing which requires the availability of zooplankton [1,2,3,4]. Yet, the most used zooplankton for the feeding of the fish larvae was Artemia [5,6]. But, the utilization, mostly in developing countries is difficult because of these cysts hatching conditions, high cost and low availability on the local market [7]. It is then important to make an intensive production of zooplankton at low cost for the expansion of fish farming. *M. macrocopa* is increasingly used as food for larval and post larval rearing of crustaceans [8] and teleost fish in culture [9,10,11,12]. It is a superior live food compared to Artemia due to its relatively high protein and nutrient content [13]. Although its culture technique is relatively simple but the specific production and feed technique knowledge for the commercial scale production was incipient in spite of its wide distribution from temperate to the tropical region.

Food resources play an important role in the production of *M. macrocopa* in natural systems [14,15]. In natural habitats, biotic and abiotic parameters such as water quality, quantity, quality level of food available and population density are one of the most important factors that interact in the population growth of the zooplankton. Among these factors, population density and food availability are the predominant factors affecting the growth of *M. macrocopa* [16]. Conventional food sources of *M. macrocopa* are very expensive and contains very low nutritive value. So, the scientists are now trying to find an alternative food source of *M. macrocopa* at low cost with high nutritive value.

40 Hence, relatively large amount of *M. macrocopa* that are required for fish larvae cultivation can be 41 produced from inexpensive, renewable waste materials. However, different types of wastes generated 42 day by day in extensive quantities, creating a significant problem in its management and disposal. 43 Besides, domestic policy of South Korea totally banned the ocean dumping of all wastes from 2014. 44 following this banning; Korean policy and industry have been tending to convert the waste into 45 resources [17]. Animal manures have a long history of use as a source of soluble phosphorous, 46 nitrogen and carbon for natural food production [18]. Animal manure used as organic matter supplied 47 to ponds can stimulate the phytoplankton growth and also increases the biomass of zooplankton 48 [19,20]. Animal wastes using for fertilization practices are popular in many countries to sustain 49 productivity at low cost [21, 22]. But, only limited information is available on the utilization prospects of 50 food waste as an alternative or additional protein source of *M. macrocopa*, which can lower the cost of 51 fish farming and at the same time, conserve the ecological value of fish ponds.

When Cladocera is used as a food for larval fish, nutrient enrichment is necessary as is the case with Artemia [23]. Because the ingredients that compose their body change according to the food they <u>consumeintakes</u> [24]. Some quantitative data are available on the fatty acid profiles of rotifers, copepods and cladocerans using algae as food, but <u>there is dearth of data no data are available</u> on the fatty acid profiles of *M. macrocopa* using organic wastes as food. *Chlorella vulgaris* is commonly used in M. macrocopa culture [41]. But it needs to be enriched by a commercial enrichment diet before feeding to the fish larvae [25]. However, it is necessary to improve the fatty acid composition of M. macrocopa in natural way by switching its diet to organic wastes. Highly unsaturated fatty acids (HUFA) enhance the essential lipid levels and these essential fatty acids promote the growth of *M.*macrocopa [26]. So, it needs to be investigated because the synthesis and accumulation of fatty acids in zooplankton are related to the stage of the individual and the frequency of reproduction among others.

The present study was designed to test the effect of different animal manures and food waste in mass culture of *M. macrocopa* with a view of investigating quality and quantity required for maximizing production. Determination <u>of the</u>-optimum concentration of each organic waste for the culture of *M. macrocopa* are important for its mass cultivation. Development of a suitable culture media for commercial production of *M. macrocopa* will be an inexpensive alternative approach to live feeds needed for fish rearing.

### 70 2. MATERIALS AND METHODS

### 71 2.1 Source of M. macrocopa

*M. macrocopa* were collected from a pond near Tongyeong, South Korea and the sample was taken to the laboratory immediately. *M. macrocopa* species were isolated from the collected sample by the micropipette and placed individually in petri dishes filled with dechlorinated tap water (10ml/plate) for breeding. Mature Moina started breeding overnight and baker's yeast was added at 1g/l to the petri dishes as a food source during the breeding period. Newborn *M. macrocopa* were collected for subsequent experiments.

### 78 2.2 Source of Organic Wastes

Three different types of animal manures such as pig manure, cow manure and poultry manure were sourced from the pig, cow and poultry production institute in Goseong, South Korea. The food wastes used in the present study included food processing waste (e.g., various types of fruit peels and leafy vegetables, rice bran, and soybean meal) and post-consumption waste (e.g., rice grain, spaghetti, beef, pork, and chicken) collected from local hotels and restaurants. The collected food wastes were transferred to the laboratory, for further processing. The food wastes were <u>mixed mixing</u> together in a

- 85 mixer machine, diced into small pieces, and excessive water was squeezed out by waste compressing
- 86 equipment. Then the final leachate was used in this experiment.

### 87 2.3 Experimental design

### 88 2.3.1 Population growth experiment

89 The experiment was conducted in total 60 tanks with a water volume of 40 L. Three replicates were 90 used for each treatment. The tanks were cleaned and dried for two days and filled with tap water and 91 left for one day with aeration for dechlorination. Water temperature in the tanks was maintained at 92 25°C. The temperature of water reservoir was regulated by a thermostat, which controlled the on / off 93 switch of a 2000-W electric heater. Four treatments with five different concentrations: chicken manure, 94 pig manure, cow manure, and food waste of 500, 1000, 2500, 5000, and 10000 ppm were assigned in 95 the experiment. Twenty healthy individuals *M. macrocopa* were individually introduced into the tanks. 96 The trial has repeated a total of three times and data pooled at the end of the period for each 97 treatment.

98 Following initiation of different growth experiments, the number of living individuals of each tank was 99 counted daily. The population of *M. macrocopa* was recorded by using the Sedgewick-Rafter counter 100 cell which is 50 mm long, 20 mm wide and 1 mm deep. *M. macrocopa* cultured in each experimental 101 tanks was recorded by using a tally counter under a dissecting microscope (10X to 40X magnification). 102 The number (no./mL) was calculated according to the formula outlined by Boyd and Lichktoppler [27] :

Number of Maine managements and -	T x 1000
Number of Mouna macrocopa /mi –	A x N x Vol. of concentrate in ml/Vol. of sample
Where,	
T = Total number of <i>M. macrocopa</i> count	ed
A = Area of grid in $mm^2$	
	Number of <i>Moina macrocopa</i> /ml = Where, T = Total number of M. macrocopa countA = Area of grid in mm2

108 1000 = Area of counting chambers in  $mm^2$ 

N = Number of grids counted

### 109 2.3.2 Water quality parameters

Dissolved oxygen (mg/L), pH and salinity(%) were measured by dipping into the water surface. Ammonia was measured by Palintest compact ammonia duo meter. Recordings were taken after tank inoculation and thereafter every 24 hours.

113 2.3.3 Vial test

114 Twenty vials of 50mL size were selected for this test and each was replicated three times. Four 115 treatments with five different concentrations: chicken manure, pig manure, cow manure, and food 116 waste of 500, 1000, 2500, 5000, and 10000 ppm were assigned in each vial. One healthy neonate of 117 less than 24hrs old was transferred in each vial and the alive individual and offspring produced from it were quantified every 24hrs. The neonates produced by M. macrocopa were collected gently and 118 119 transferred into a culture dish for quantification. This counting was also carried out using a tally counter. 120 Then it transferred to the new test jars with appropriate culture medium and the dead adults and 121 neonates were removed. Each vial test was discontinued after the last adult in each vial was died. 122 Mortality and fecundity were recorded to calculate the life table demography of *M. macrocopa*.

### 123 2.3.4 Life table demography

Life table demographics is an important tool for describing the life cycle of zooplankton under continuously changing environmental conditions. The survival period, initial age of reproduction, average longevity, gross reproduction rate, net reproduction rate, rate of increase, and generation time were selected for life history variables for this study [28]. The following definitions apply: initial age of reproduction = the time when a female started to produce her first batch of offspring (number of days); longevity = the average number of days the female survived. The following formulae were used [29].

130 Average longevity =  $\sum n_{\chi}/n$ ;

- 131 Gross Reproduction rate =  $\sum m_{\chi}$ ;
- 132 Net reproduction rate (Ro)= $\sum I_{\chi} m_{\chi}$ ;
- 133 Generation time (T) =  $\sum I_{\chi} m_{\chi} \chi / Ro$
- 134 Where,

- 135 n x =Number of individuals alive for each age class
- 136 m=The age specific fecundity (number of neonates produced per surviving female at age X)
- 137 I  $_{\chi}$  =The proportion of individuals surviving to age  $\chi$
- 138 n = The number of replicates

The final rate of population increase (r) was calculated over the 21 days experimental period. The rate of population increase (r) was derived using the following equation;  $r = (InN_t-InN_0)/t$ , where N<sub>0</sub>=initial population density and N<sub>1</sub>=population density after time t [29].

### 142 **2.3.5 Population density of M. macrocopa**

Twenty vials of 50 mL size were assigned with four treatments of five different concentrations: chicken manure, pig manure, cow manure, and food waste of 500; 1,000; 2,500; 5,000; and 10,000 ppm in each vial. One healthy neonate of less than 24hrs old was transferred in each vial and the alive individual and offspring produced from it were quantified every 24hrs, which was carried out until finishing of this experiment. *M. macrocopa* were transferred into different culture dishes for quantification and after quantification, live *M. macrocopa* were returned to the culture vial, and the dead organisms were discarded. This experiment was carried out for 21 days.

### 150 2.3.6 Fatty acid analysis of M. macrocopa cultured in different organic wastes

151 Total lipids of *M. macrocopa* were extracted according to the Bligh and Dyer method [30] by using 152 solvent mixture consisting of chloroform and methanol (2:1, v/v). After phase equilibration, the lower 153 chloroform layer was removed and total lipids were extracted by removing solvent using a rotary 154 evaporator (R-114, BUCHI, Swiss) at 38 °C. 100 mg of extracted total lipid were put into a capped tube 155 and added 1.5 ml 0.5 N NaOH-methanol solution. The sample was mixed by vortex and heated 100 °C 156 for 8 minutes for saponification. After cooling, methylation was done by using a fatty acid methyl ester 157 (FAME) with BF3-methanol. Then the sample was dissolved into 2 ml iso-octane and fatty acids were 158 analyzed using gas chromatography (Clarus 600, Perkin Elmer, USA) equipped with capillary column 159 (Omegawax-320, 30 m × 0.25 mm I.D., Supelco Co., Bellefonte, PA, USA). The operating parameters 160 were as follows: carrier gas =helium; detector (FID) temperature =270°C; injection temperature = 161 250 °C; column temperature =180 °C for 8 min, programmed to increase at 3 °C/min up to 230 °C with a

162 final holding time of 10 min; split injection at 1:50 ratio. Menhaden oil was used as standard. Each of 163 the specific fatty acid methyl ester peaks was identified by determining its equivalent chain length with 164 reference to the known standard.

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### 166 2.4 Statistical Analysis

The statistical analysis were carried out to evaluate the differences in the means of the derived individual number of *M. macrocopa* and environmental parameters of different treatments by using one-way ANOVA. Statistical significance among the different treatments was accepted at p<0.05 and the statistical package of SPSS- 16 (SYSTA, USA) was used to express the result.

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### 172 **3. RESULTS**

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### 174 **3.1 Water quality**

175 Table 1 shows the mean pH, DO and ammonia content of five various concentrations of four different 176 culture medium over three weeks experimental period. The highest pH was recorded in the pig manure 177 treatment which was in the range of between 7.33 to 7.72 and the lowest was recorded in the food waste which was in the range of 5.50 to 6.46. pH increased with the increase concentration of animal 178 179 manures, but in case of food waste it decreased. Food waste treated media showed significantly 180 lowest (P<0.05) DO level throughout the culture period, which were found to be in the range of 0.4 to 181 1.08 mg/l. As the temperature was fixed from the beginning of the experiment, there is no significant 182 difference was observed between the treatments. Ammonia contents increased with the increasing 183 concentrations of organic wastes but showed no significant difference (P >0.05) among the treatments 184 of all concentration.

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# Table 1. Water quality parameters for the M. macrocopa cultures at different concentrations of animal

manures and food waste of the experimental duration

Culture	Concentration	pН	DO	Temperature	Ammonia	Salinity
medium						

	(ppm)		(mg/l)	(°C)	(mg/l)	(PSU)
	500	6.816±0.02	2.83±0.01	25±0.1	0.20±0.03	0.01±0.0
	1000	6.973±0.02	2.71±0.01	25±0.1	0.22±0.03	0.01±0.0
Chicken manure	25000	7.281±0.04	2.25±0.02	25±0.1	0.33±0.06	0.01±0.0
	5000	7.406±0.03	2.28±0.03	25±0.1	0.39±0.04	0.02±0.1
	10000	7.554±0.02	0.57±0.03	25±0.1	0.41±0.08	0.02±0.0
	500	7.335±0.01	3.54±0.02	25±0.1	0.23±0.02	0.01±0.0
	1000	7.208±0.02	3.23±0.02	25±0.1	0.25±0.06	0.01±0.0
Pig manure	25000	7.557±0.02	3.06±0.04	25±0.1	0.29±0.05	0.01±0.0
	5000	$7.524 \pm 0.03$	2.86±0.07	25±0.1	0.41±0.03	0.02±0.0
	10000	7.729±0.06	1.03±0.03	25±0.1	0.44±0.05	0.03±0.0
	500	6.719±0.02	3.45±0.01	25±0.1	0.19±0.05	0.01±0.0
	1000	7.083±0.01	3.11±0.03	25±0.1	0.22±0.04	0.01±0.0
Cow manure	25000	7.159±0.03	1.76±0.02	25±0.1	0.26±0.04	0.01±0.0
	5000	7.230±0.02	1.53±0.05	25±0.1	0.36±0.03	0.01±0.0
	10000	7.592±0.04	0.61±0.03	25±0.1	0.37±0.08	0.03±0.1
	500	6.461±0.02	1.08±0.01	25±0.1	0.23±0.02	0.01±0.0
	1000	6.823±0.03	0.91±0.02	25±0.1	0.25±0.04	0.01±0.0
Food waste	25000	6.064±0.05	0.93±0.01	25±0.1	0.36±0.06	0.01±0.0
	5000	5.963±0.02	0.32±0.01	25±0.5	0.35±0.05	0.03±0.1
	10000	5.501±0.03	0.41±0.02	25±0.1	0.38±0.04	0.03±0.0

### 190 **3.2 Population growth of M. macrocopa**

191 Figure 1 shows the population growth of *M. macrocopa* with five various concentrations of four 192 different organic waste treated medium for 21 days experimental period. Population growth was 193 significantly higher in the treatments with low concentrations, on the contrary there is no population 194 growth was observed in 10,000 ppm; the highest concentration used in this experiment. The highest 195 growth was recorded in 1,000 ppm concentration for four culture medium and among these four 196 mediums, food waste showed the best growth rate. Figure 2 showed the mean population growth of M. 197 macrocopa cultivated in 1,000 ppm concentration of four culture medium. Highest mean population 198 was found in food waste medium which was followed by chicken manure medium and showed a 199 significant difference (P<0.05) among the other medium, where there is no significant (P<0.05) 200 difference was observed between pig manure and cow manure treated treatment.



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Fig. 1. Population growth of *M. macrocopa* in different concentrations of (A) chicken manure, (B) pig manure, (C) cow manure, & (D) food waste culture medium for 21 days experimental period. Error bars indicate means ± standard deviation.



Fig. 2. Mean population growth of *M. macrocopa* cultured in 1000ppm concentration of four culture medium. Values are the (mean  $\pm$  SD). Different subscripts denote significant differences at P < 0.05 (a < b < c).

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## 211 **3.3 Life table demography**

212 Data on the selected life history variables (Table 2) of *M. macrocopa* showed that the average lifespan 213 was lowest in treatment with high concentration, about 1 to 2 days. The offspring production of M. 214 macrocopa in relation to the different concentrations of treatments showed a distinct shift towards early 215 reproduction with low concentration treatments. Gross and net reproductive rate also showed similar 216 trends. Large numbers of offspring were produced by M. macrocopa cultured in food waste (500; 217 1000ppm), chicken manure (500; 1000 ppm/l), cow manure (500; 1,000 ppm) and pig manure (500, 218 1,000 ppm) medium. Fecundity declined at high concentrations (2500 ppm and 5000 ppm), while the 219 highest 10,000 ppm concentration did not lead to the production of offspring for every treatments. The 220 rate of population increase was positive for all the culture medium, but 1000 ppm/l showed the 221 maximum. The highest rate of population increase (r) calculated for this experiment was 0.51 ± 0.08 222 obtained in the food waste treated treatment. This was followed by the chicken manure, cow manure 223 and pig manure medium with 'r' value of 0.47±0.23, 0.43±0.16, and 0.33±0.02 respectively.

# 225Table 2. Life table of *M. macrocopa*, cultured with different organic wastes at different226concentrations. Data are the means with the standard error of three replicates

_	Medium types	Medium Conc. (ppm)	Initial age of reprod- uction	Longevity	Net reprod- uction rate	Gross repr- oduction rate	Generation time	Rate of populati -ion increase
		500	2.89±0.06	8.16±0.13	11.73±0.11	14.06±1.03	3.41±0.11	0.39
	Chicken	1000	2.87±0.08	9.20±0.23	19.32±0.08	23.42±0.91	3.00±0.36	0.47
	manure	2500	3.00±0.08	6.27±0.18	9.17±0.13	11.42±0.85	3.16±0.25	0.36
		5000	3.07±0.05	5.63±0.20	5.21±0.16	5.21±1.14	3.27±0.25	0.32
		10000	-	2.86±0.41	-			
_		500	3.06±0.03	5.63±0.13	8.25±0.21	10.14±0.21	4.00±0.17	0.24
	Pig	1000	3.08±0.08	7.74±0.11	11.69±0.17	13.32±0.16	3.53±0.26	0.33
	manure	2500	3.00±0.07	6.75±0.18	7.32±0.17	9.11±0.20	3.60±0.25	0.27
		5000	-	1.00±0.20	7.	· ·	-	-
		10000	- 🦿	$1.00 \pm 0.41$		-	-	-
		500	3.06±0.08	7.63±0.11	10.21±0.21	10.53±1.11	3.23±0.11	0.33
	Cow	1000	3.06±0.05	9.14±0.16	15.57±0.18	17.71±1.31	3.10±0.36	0.43
	manure	2500	3.26±0.08	9.17±0.28	15.42±0.16	17.24±0.93	3.00±0.25	0.43
		5000	3.33±0.05	5.00±1.22	4.16 ±0.16	4.23±1.14	3.27±0.25	0.32
•		10000		3.75±0.41	-	-	-	-
-		500	3.00±0.07	8.71±0.11	12.42±0.18	15.35±1.00	3.00±0.10	0.43
		1000	2.88±0.06	9.82±0.26	28.16±0.18	33.71±0.21	3.00±0.16	0.51
	Food	2500	-	1.75±0.08	-	-	-	-
	waste	5000	-	1.00±0.01	-	-	-	-
		10000	-	1.00±0.00	-	-	-	-

Dash (-) indicates no offspring was produced

### 228 3.4 Populations density of M. macrocopa

229 Populations from a single neonate of *M. macrocopa* showed average growth rate until the first week in 230 various concentrations of four different culture medium (Figure 3). It was growing continuously until the 231 second week and after that, it started to decline and this same trend was observed in all treatments. 232 Among all the concentrations highest population was found in 1,000 ppm concentration of all the 233 culture medium. In case of food waste medium neonate is died in 2500, 5000 and 10000 ppm 234 concentrations within 24hrs and 75 numbers of individual M. macrocopa were counted on day 10, in 235 1,000 ppm concentration, which was its peak population. Vials containing chicken manure, cow 236 manure and pig manure medium showed moderate production with a peak population of 57, 52 and 43 237 individual on day 9 in 1,000 ppm concentration. After that the decreasing trend was started, which 238 means that these peak populations are the maximum density of M. macrocopa for different culture 239 medium.



Fig. 3. Population density from a single neonate of *M. macrocopa* in different concentrations of (A) chicken manure, (B) pig manure, (C) cow manure, & (D) food waste treated medium for 21 days experimental period. Error bars indicate means ± standard deviation.

### 245 **3.5 Fatty acid composition of M. macrocopa cultured in different organic wastes**

Table 3 shows the average percentage of fatty acid composition of *M. macrocopa* cultured in different organic waste. Among the saturated fatty acids, 14:0, 16:0 and 18:0 comprised of about 33% of the total fatty acids and 16:1n-7, 18:1n-7, 18:2n-6 are the most dominant unsaturated fatty acids. The fatty acid composition of *M. macrocopa* cultured in cow manure, food waste, chicken manure and pig manure was dominated by, 16:00, 18:00, 18:1n-9, 18:2n-6. The level of EPA and DHA was significantly higher in *M. macrocopa* that cultured in food waste medium than those other wastes.

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### 253 4. DISCUSSION

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255 Among the cladocereans, M. macrocopa has been investigated most intensively with regard to the 256 effects of food abundance on its growth and reproduction [10,31,32]. Quality and quantity of food are 257 the most important factors in determining biomass production of *M. macrocopa* species. The results of 258 this study indicate that the growth performance of M. macrocopa using various concentrations of 259 different organic wastes as a culture medium are different. Low population growth was observed in 260 high concentration of culture medium, while the highest concentration used in this study did not lead to 261 the production of offspring in all the culture medium. This phenomenon has been attributed by various 262 workers that the presence of high concentrations of animal manure significantly reduced the water 263 quality, deplete the plankton population and cause adverse effect on the culture [4]. Nandini and Sarma [33], revealed that, the decline in neonate production that accompanied increasing 264 265 concentrations of culture medium were presumably caused by the increased effort associated with 266 food gathering due to active filtering of the food particles. In fact, high concentrations of all the diets 267 produced suboptimal culture conditions. Burak [34] & Porter[35] described that, high concentration of 268 particles can actually lead to starvation of cladocerans as they are unable to clean thoracic limbs that 269 are clogged by high particulate concentrations. Savas [31] also found that, population of *M. macrocopa* 270 declined in using high concentration of algal supplement. In this study, 1000 ppm concentration 271 showed the optimal concentration of all the culture media in terms of growth and reproduction

waste :0.18 ±2.71 ±0.09 ±0.26 ±0.28 0.90
:0.18 ±2.71 ±0.09 ±0.26 ±0.28 0.90
±2.71 ±0.09 ±0.26 ±0.28 0.90
±0.09 ±0.26 ±0.28 0.90
±0.26 ±0.28 0.90
±0.28 0.90
0.90
±1.25
:0.14
-
:0.07
±0.05
±0.09
±0.04
±0.06
:0.07
±0.01
±0.01
.75
.57
43

#### Table 3. Fatty acids composition (%) of *M. macrocopa* cultured in different organic wastes

Data of 1,000 ppm concentration of different organic waste are shown here. Hyphen (-) indicates non- detectable fatty acids.

279 performance. Among the culture medium, the highest population growth was observed in food waste 280 medium. In this study pH increased with the high concentration of animal manures but food waste 281 showed different trend where pH decreased with the increasing concentrations, might be the presence 282 of higher amount longer chain fatty acids. Life table demography of *M. macrocopa* followed the same 283 trend. Average lifespan was lowest in culture mediums with higher concentration. Food waste 284 containing 1,000 ppm concentration showed the highest average lifespan and early reproduction ability. 285 The present study showed that, *M. macrocopa* needs time to became sexually mature at high diet 286 concentration. In contrast, Loh [32] reported that, initial age of reproduction of *M. macrocopa* is earlier 287 in high concentration diet than in low concentration. Different results are observed in this test, which indicates that diet type and concentration play a significant role in determining the initial age of 288 289 reproduction. Gross and net reproduction rates were generally higher at lower treatment 290 concentrations and highest in food waste medium than other diets. Jana and Pal [36] revealed in their 291 study, high fecundity and gross reproduction rates suggest that growth performance of any species is 292 largely depends on the high carbon/nitrogen ratio in the food source. Which clearly indicates that food 293 waste contained high C/N ratio than the other medium.

294 The highest population density was obtained in 1,000 ppm concentration of food waste medium. The 295 good performance of this food source can be attributed to the feeding habit of *M. macrocopa* that tend 296 to consume bacteria and filtered particles that are abundant in food waste, when other food sources 297 has limited. In terms of time and efficiency, M. macrocopa cultured in food waste reached its peak 298 population on day 10 which was about 75 individual. This also means that M. macrocopa could be 299 harvested by food waste within a shorter period of time, thus allowing more number of cultivation 300 batches per cycle which is important for commercial live feed producers. After reaching the peak 301 population day within second week, the population began to decrease from the starting of third week. 302 Which could be caused by insufficient of space, food availability, sexual transformation, and/or 303 allelopathic effects [37,10,38,39,40]. Jiun [41] reported that, M. macrocopa has a higher density 304 adaptation in a captive culture environment compared to M. micrura, because high stocking density 305 may possibly lead to a population collapse. According to Jana and Pal [36], the growth performance of 306 M. macrocopa was limited at the density of 4 ind. and 20 ind./ 50ml. Results of this study showed that, 307 M. macrocopa had a better adaptation in food waste treatment at the density up to 75 individual per 308 50mL which is more higher than that previous study.

309 Muller et al., [26] revealed that, population growth and reproduction of the species depends on the 310 quantity of reserve lipids (14:0, 16:0 and 18:0). M. macrocopa cultured in this study comprised of 33% 311 of saturated fatty acids in each treatment. M. macrocopa exhibited a fatty acid profile of poly 312 unsaturated fatty acids (18:1n-9, 18:2n-6 and 18:3n-3) constituting 45.1%, 37.6%, 36.4% and 34.3% 313 when grown in cow manure, food waste, pig manure and chicken manure treatment respectively. It 314 could be that a decrease in PUFA, reduces the capacity of animals to withstand in the environment as reflected in adverse changes in demography [42]. Comparatively higher percentage of EPA and DHA 315 316 was found in *M. macrocopa* grown in food waste treatment and the levels of EPA (1.8%) were 317 comparable with those in rotifers or Artemia that were fed algal diets in another study [10]. This result 318 suggests that *M. macrocopa*, when culture in food waste treatment, has the potential to be a suitable 319 diet containing essential n-3 HUFAs for fish larvae.

Our study demonstrated that *M. macrocopa* can be cultured using animal manure and food waste. This is also in agreement with Nandini [33] and Golder [43]. However, Siebe C [44] reported that, Moina sp. cultivation using domestic wastes as a food source posing a high risk of pathogen contamination or toxicant pollution. But, in case of food waste this contamination possibilities is comparatively low. Studies has shown that food waste can replace part of the fish meal used in fish feeds to produce quality fish and no health risk was observed in the health risk assessment test [45].

### 326 5. CONCLUSION

In conclusion, the results of this experiment suggest that *M. macrocopa* could be cultivated using number of the concentration of different animal manure and food waste. Results indicate that, the food waste appeared to be more effective compared to all other treatments for successful mass culture of *M. macrocopa* to high density and higher percentage of n-3 HUFA, which may serve as effective, inexpensive and sustainable food sources for *M. macrocopa* cultivation.

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### 335 COMPETING INTERESTS

The authors declare that they have no competing interests.

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### 338 ETHICS APPROVAL

339 All experimental protocols followed the guidelines of the Institutional animal Care and Use Committee

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340 of the Gyeongsang National University.

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