

A comparison between marine and terrestrial invertebrate meals for mirror carp (*Cyprinus carpio*) diets: impact on growth, haematology and health

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DOI: 10.1111/are.13318



Wan, A.H.L., Snellgrove, D.L. and Davies, S.J. 2017. A comparison between marine and terrestrial invertebrate meals for mirror carp (*Cyprinus carpio*) diets: Impact on growth, haematology and health. *Aquaculture Research*.

7 April 2017

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21 **Abstract**

22 Invertebrate-meals (e.g. polychaetes and insects) present novel and sustainable high
23 quality nutrient sources for use in fish feed formulations. To test this innovative source,
24 an eleven-week feeding trial was conducted evaluating the effects of replacing the
25 fishmeal (FM) component as an example of a superior protein source (FM CTRL) with
26 ragworm meal (RW, *Nereis virens*), or/and silkworm pupae (SWP, *Bombyx mori*) in
27 mirror carp (*Cyprinus carpio*) diets. Three experimental diets with partial replacement
28 of FM (diets: RW+FM, SWP+FM, and RW+SWP+FM) were formulated. All diets were
29 formulated to be *iso*-nitrogenous, *iso*-lipidic, and *iso*-energetic. Growth performance
30 and feed utilisation indices were assessed, and the feeding trial concluded with the
31 analysis of haematological parameters to provide an indication of carp physiological
32 and health status. Mean weight gain was greatest in mirror carp fed RW+FM (60.83
33 fish⁻¹ day⁻¹; $P < 0.05$ vs all other diets) followed by SWP+FM (40.62 g fish⁻¹ day⁻¹;
34 $P < 0.05$ vs all other diets). The least weight gain was achieved in fish fed
35 FM+SWP+RW+ and FM CTRL (34.34 g fish⁻¹ day⁻¹ and 33.96 g fish⁻¹ day⁻¹,
36 respectively; not significantly different from each other). Fish fed on RW+FM diet had
37 significantly lower plasma ammonia concentrations than any other dietary groups
38 ($P = 0.04$). Mirror carp fed on SWP+FM diet (111.52 units mL⁻¹) were observed to have
39 a marked enhancement in alternative complement activity than FM CTRL (79.21 units
40 mL⁻¹, $P = 0.041$). Both ragworm and silkworm pupae meal present attractive sustainable
41 functional feed component in carp diets, with benefits on enhancing growth
42 performance and specific physiological parameters.

43

44 **Keywords:** Invertebrate meal; Carp; Silkworm pupae; *Bombyx mori*; Ragworm; *Nereis*
45 *virens*.

46 **Introduction**

47 Finfish aquaculture has expanded to become one of the largest and fastest sectors in the
48 food production industry. In 2013, 70.5 million tonnes of farmed food fish were
49 produced, and since 1980 the production level of world aquaculture has increased at an
50 annual rate of 8.6% (FAO, 2014). Consequently, the demand for aquafeeds and its main
51 protein constituent, fishmeal, has also dramatically increased. In 2009, it was estimated
52 that 30% of capture fisheries production was processed into fishmeal and fish oil (Olsen
53 & Hasan, 2012). However, with dwindling wild fish stocks, fishmeal is now considered
54 to be a finite protein source and to be used strategically. Responding to the bottleneck in
55 aquaculture production, many governments, researchers and aquafeed manufacturers
56 have now sought to evaluate possible alternatives that are deemed more sustainable.

57

58 Previous studies on fishmeal alternatives include various fisheries by-products (Toppe,
59 Aksnes, Hope, & Albrektsen 2006; Lee, Powell, Barrows, Smiley, Bechtel & Hardy
60 2010), terrestrial animal by-products (Davies, Gouveia, Laporte, Woodgate & Nates
61 2009), single-cell organisms (Lunger, Craig & McLean 2006; Zerai, Fitzsimmons,
62 Collier & Duff 2008), algae (Soler-villa, Coughlan, Guiry & Kraan 2009; Xu, Zhang,
63 Wu, Liu, Wang, You & Li 2011) and plant meals (Opstvedt, Aksnes, Hope & Pike
64 2003; Torstensen, Espe, Sanden, Stubhaug, Waagbø, Hemre, Fontanillas, Nordgarden,
65 Hevrøy, Olsvik & Berntssen 2008). However, in some instances fishmeal replacement
66 candidates can cause a variety of physiological problems in the fish, e.g. depressed
67 growth rates, nutritional health problems or reduced palatability (Deng, Mai, Qinghui,
68 Zhang, Wang, Xu, Liufu 2006; Hardy, 2010). These are often attributed to the presence
69 of Anti-Nutritional Factors (ANF's), reduced nutrient availability, and/or the
70 deficiencies in essential nutrient(s), e.g. amino acid, vitamins and trace metals (Francis,

71 Makkar & Becker 2001; Opstvedt *et al.* 2003; Fasakin, Serwata & Davies 2005; Hansen
72 Rosenlund, Karlsen, Wolfgang & Gro-Ingunn 2007).

73

74 In recent years, the use of invertebrate meals has gained much interest as a sustainable
75 alternative to fishmeal (Barroso, De Haro, Sánchez-Muros, Venegas, Martínez-Sánchez
76 & Pérez-Bañón 2014; Henry, Gasco, Piccolo, & Fountoulaki 2015). Invertebrate meals,
77 such as black soldier fly, housefly, and silkworm pupae have been tested in a range of
78 dietary inclusion levels (5 to 30%) in African catfish (Ng, Liew, Ang & Wong 2001),
79 channel catfish (Bondari & Sheppard, 1987), chum salmon (Akiyama, Murai, Hirasawa
80 & Nose 1984), common carp (Nandeesh, Gangadhara & Manissery 1999), rainbow
81 trout (St-Hilaire, Sheppard, Tomberlin, Irving, Newton, McGuire, Mosley, Hardy &
82 Sealey 2007), and blue tilapia (Bondari & Sheppard, 1987). These studies have shown
83 that invertebrate meals have promising attributes, providing a sustainable protein rich
84 alternative, an adequate amino acid profile, highly digestibility, and improving fish
85 growth performance (Barroso, De Haro, Sánchez-Muros, Venegas, Martínez-Sánchez,
86 & Pérez-Bañón, 2014).

87

88 Ragworm (*Nereis virens*) is a marine polychaete worm found in the benthic strata of
89 northern hemisphere estuarine habitats. Preliminary studies have indicated its suitability
90 as a feed ingredient for both crustacean and finfish production (Day, Howell & Jones
91 1997, Salze, McLean, Battle, Schwarz & Craig 2010). Similarly, meals derived from the
92 silkworm (*Bombyx mori*) pupae have been well documented as suitable fishmeal
93 alternatives in diets of meat-producing terrestrial species (Ijaiya & Eko, 2009; Medhi,
94 Nath, Gohain, & Bhuyan 2009). Growth studies performed on carp species (Nandeesh,
95 Gangadhara, Varghese & Keshavanath 2000; Rangacharyulu, Giri, Paul, Yashoda, Rao,

96 Mahendrakar, Mohanty & Mukhopadhyay 2003) and walking catfish (*Clarias*
97 *batrachus*, Habib, Hasan, Akand & Siddiqua 1994) have shown that replacement of
98 fishmeal by silkworm pupae meal maintained or exceeded the growth performances
99 compared to the dietary reference treatment group. More recently however, Ji, Zhang,
100 Huang, Cheng & Liu (2015) tested the feasibility of replacing fishmeal with silkworm
101 meal for juvenile Jian carp in multi-ingredient formulated diets containing just 10%
102 fishmeal. The study reported reduced growth rate and impaired anti-oxidant enzyme
103 status, decreased digestive function and unfavourable changes in hepatic and intestinal
104 morphology for these carp fed higher silk worm pupae meal dietary inclusion.

105

106 Cyprinids (carp) are an important freshwater farmed fish in Eastern Europe, India and
107 China where demand is high. In 2010, China has produced over 15 million tonnes of
108 farmed cyprinids, ranging from grass carp (*Ctenopharyngodon idella*) to common carp
109 (*Cyprinus carpio*) (Chiu, Li, Guo, Bai, Fedor & Naylor 2013). Furthermore, Koi are the
110 genetic variant of the domesticated farm carp, and are highly prized in the ornamental
111 pet fish trade as companion animals (FAO, 2014) in many parts of the world. It is
112 therefore important to assess the potential of novel feed ingredients for this species in
113 the context of sustainability with respect to the needs of both the aquaculture and
114 aquatics industries. For this reason, an investigation was directed to assess a
115 commercially produced marine invertebrate (polychaete worm meal) and a
116 commercially produced terrestrial invertebrate (silkworm meal) tested singly and in a
117 combination, in experimental diets for mirror carp substituting the fishmeal component
118 in formulated experimental diets for mirror carp.

119

120 **Materials and Methods**

121 The present study examined the effects on both growth performance and haematological
122 parameters (standard, biochemical and immunological), when ragworm or/and
123 silkworm pupae meal was assessed against a single primary protein (fishmeal) in
124 balanced diets for juvenile mirror carp (*Cyprinus carpio*) as a standard protein of
125 defined Biological Value (BV).

126

127 **Fish and experimental facilities**

128 An eleven week feeding trial was conducted at the Aquaculture and Fish Nutrition
129 Research Aquarium, University of Plymouth. Mix-sex mirror carp ($n=240$) were
130 sourced from Hampshire Carp Hatcheries (Bowlake fish farm, UK) and acclimatised for
131 5 weeks in an experimental re-circulated system. During acclimatisation fish were fed
132 on a commercial EWOS micro 50P diet (EWOS Ltd., UK). The experimental re-
133 circulated system comprised 12 (38 cm x 38 cm x 50 cm) fibreglass tanks suspended
134 over a 900 L filter sump tank. The core experimental design consisted of four test diets
135 each with three replicate test groups randomized over the twelve tanks in the holding
136 system. Fish were graded and stocked at a density of 20 fish per tank, with an initial
137 mean fish weight of 14.9 ± 0.13 g (\pm S.E.M, $n=20$). Each tank had an aerated flow rate
138 of 4 L min^{-1} and was maintained at 24.6 ± 0.2 °C with a dissolved oxygen level of $>89\%$.
139 Nitrogenous waste levels were monitored weekly and maintained to values of (means \leq ,
140 mg L^{-1} , \pm S.E., $n= 8$) ammonium, 0.1 ± 0.02 ; nitrite 0.03 ± 0.01 and nitrate 38 ± 6 [Hach
141 Lange, Salford, UK]. Photoperiod was set on a diurnal cycle of 12 h light and 12 h
142 darkness. Diets were given at a ration level of 3% of body weight per day. Batch group

143 weights of each tank were measured on a weekly basis to determine the growth
144 performance:
145 Weight gain: Final weight [g] - Initial weight [g]
146 Specific growth rate (SGR): $\ln(\text{final body weight [g]}) - \ln(\text{initial body weight [g]}) / \text{time}$
147 [days] x 100
148 Food conversion ratio (FCR): feed fed [g] / live weight gain [g]
149 Protein efficiency ratio (PER): weight gain [g] / protein ingested [g].
150 At the conclusion of the feeding experiment, six fish from each tank were randomly
151 sampled for blood analysis that included basic haematology, biochemical and
152 immunological haematology parameters. All work was conducted according to the 1986
153 Animal Scientific Procedures Act (UK Home Office) regulations of the UK, the
154 University of Plymouth's ethical approval process, and the Ethics Committee of the
155 WALTHAM Centre for Pet Nutrition.

156

157 **Diets**

158 Three experimental diets were formulated with a protein replacement level of 6.33% for
159 silkworm pupae meal (SWP+FM diet), 7.52% ragworm meal (RW+FM diet), 3.30%
160 and 4.06% of silkworm pupae and ragworm meal, respectively (SWP+RW+FM diet).
161 Calcium chloride (2.98%) was added to SWP+RW+FM diet to maintain the calcium
162 content due to low calcium levels in this specific diet mixture compared to the reference
163 diet. This reference diet was produced with no invertebrate inclusion and providing a
164 superior basal protein BV consisting of LT (Low Temperature) fishmeal. The
165 formulation and preparation process included the use of a commercial food processor
166 [HL1400-10STDA, Hobart Food Equipment, Australia] to blend the feed materials into
167 a dough consistency and cold extruded through a PTM P6 feed extruder system

168 [Plymouth Marine Ltd., UK]. A final pellet size of 2 mm in diameter and 5 mm in
169 length was produced. Diets were dried in a dehumidifying oven at 46 °C for 48 h.
170

171 **Proximate analysis**

172 Moisture, crude protein, crude lipid, and ash in feed ingredients (Table 1) and finished
173 diets (Table 2) were determined by following AOAC (1995) standard methods. Briefly,
174 moisture was determined by heating samples to 105 °C, until a constant weight was
175 achieved. Crude protein was calculated from the nitrogen content ($N \times 6.25$) using the
176 Kjeldahl method. Samples were first acid digested using a Kjeldahltherm microsystem
177 40 [C. Gerhardt GmbH & Co. KG, Germany] and distilled in a Vapodest 40 [C.
178 Gerhardt GmbH & Co. KG, Königswinter, Germany]. Crude lipid content in samples
179 was determined through the petroleum ether extraction method, using a Soxtec extractor
180 HT 1043 extraction unit [Foss Tecator AB, Hoganas, Sweden], while crude ash content
181 was determined by igniting samples at 600 °C for 16 h and weighing the residual ash.
182 Energy values were measured using a bomb calorimeter Parr 1356 [Parr Instrument
183 Company, Illinois, US].

184

185 **Amino acid analysis**

186 Amino acid composition (Table 2) was performed on hydrolysed weighed samples of
187 feed in sealed ampoules in *vacuo* with 6N HCl at 110°C for 22 hrs. Excess acid was
188 removed by a flash evaporator under reduced pressure at below 40°C. Samples were
189 reconstituted in buffer and amino acid analysis conducted on a Biochrome 30 Amino
190 Acid Analyser [Biochrom, Cambridge, UK]. Methionine was determined separately
191 using a performic acid oxidation. Tryptophan was recovered in a separate alkaline

192 hydrolysis step according to standard protocols (Longvah, Mangthya & Ramulu, 2011;
193 Davies & Gouveia, 2000).

194

195 **Blood sample collection**

196 At the conclusion of the experiment, blood samples were collected from the
197 experimental fish to examine possible modifications in the components of carp blood,
198 which could give indications on physiological health and nutritional status (Kaushik &
199 Seiliez, 2010). Samples were collected by lightly anaesthetising fish in an aerated
200 anaesthetic bath (120 mg L^{-1} , tricane methane sulphonate, Pharmaq, Fordingbridge,
201 UK). Six fish from each tank were sampled for blood at the caudal ventral vein, using a
202 25-gauge needle and 1 mL syringe. Needles and syringes used for plasma collection
203 ($n=3$) were heparinised prior to collection (Walencik & Witeska, 2007). Blood samples
204 were immediately centrifuged at 13,000 rpm for 11 mins and collected supernatant was
205 subjected to a further 1 min centrifuge at 13,000 rpm and aliquot for storage at $-80 \text{ }^{\circ}\text{C}$.
206 For serum collection ($n=3$), blood was allowed to clot for 24 h at $2-4 \text{ }^{\circ}\text{C}$ and centrifuged
207 at 13,000 rpm for 11 mins. Supernatant was centrifuged for a further 1 minute. Samples
208 were stored at $-80 \text{ }^{\circ}\text{C}$ for later analysis.

209

210 **Basic haematology parameters**

211 Freshly collected whole blood was immediately used in the following haematological
212 parameters. Pack cell volume was determined using a micro hematocrit method, as
213 described by Dacie & Lewis (1984). Haemoglobin determination was performed
214 through the cyanmethaemoglobin method (Dacie & Lewis, 1984) using a commercial
215 Drabkin's cyanide-ferricyanide solution [Sigma Alderich, Poole, UK]. For total

216 erythrocyte, 20 μL of whole blood was fixed in 1 mL of Dacies fluid (Dacie & Lewis,
217 1984) and counts carried out by the method described in Handy & Depledge (1999).
218

219 **Haematological and related immune parameters**

220 Serum haemolytic complement activity through the alternative pathway was performed
221 as described by Yano (1992), using washed rabbit red blood cells (2×10^8 cells mL^{-1} ,
222 RaRBC) as the target cells [TCS, Botolph Claydon, England]. Briefly, the three serum
223 samples from the same tank were pooled together (250 μL) and diluted 20 fold using
224 Mg^{2+} -EGTA-GVB buffer (20 mL of 4.15 g NaCl, 0.51 g $\text{C}_4\text{H}_3\text{N}_2\text{NaO}_3$, 1.75 mL 1 N
225 HCl, dissolved in 100 mL dH_2O , pH 7.5; 10mL of 3.8 g EGTA, 2.03 g magnesium
226 chloride hexahydrate, 0.7 g NaOH, dissolved in dH_2O , pH7.5; 70 mL of 0.1 g gelatin,
227 dH_2O , pH 7.5). Diluted serum was mixed with 100 μL of RaRBC suspension and
228 incubated for 90 mins (20 $^\circ\text{C}$), with occasional shaking. The reaction was ceased by
229 adding 3.15 mL cold phosphate buffer saline to the suspension and centrifuged at
230 10,000 rpm for 5 min. Supernatant was measured for its maximal absorbance at 414 nm.
231 Total haemolysis was carried out by adding 100 μL RaRBC suspension to 3.4 mL
232 dH_2O . The extent of haemolysis for each serial dilution was calculated by following the
233 equation: Haemolysis (ACH_{50}) = $1/k \times (\text{reciprocal of initial dilution}) \times 0.5$.

234

235 Serum lysozyme activity determination was achieved using the turbidimetric method as
236 described by Ellis (1990). Briefly, 50 μL of serum sample was mixed into 1.95 mL
237 suspension of 0.2 mg mL^{-1} *Micrococcus lysodeikticus* [Sigma Alderich, Poole, UK] in
238 0.05 M sodium phosphate buffer (pH 6.2). Reduction of turbidity was measured (450
239 nm) after mixing from time 0 to 5 mins at 20 $^\circ\text{C}$. Lysozyme activity is defined as one
240 unit of enzyme producing an absorbance decrease of 0.001 min^{-1} .

241 **Haematological biochemistry**

242 Total serum protein was assayed on a 96 multi-well plate, using commercial Bradford's
243 dye solution [Sigma Aldrich, Poole, UK]. Four microlitres of diluted serum sample
244 (1:40) was mixed with 200 μ L of Bradford solution. The mixture was agitated for 30
245 seconds before incubation at 25 °C for 10 mins and was read on a microplate reader
246 (595 nm). Albumin was determined using a bromocresol green dye- binding method
247 (Spencer & Price, 1977). Serum samples (20 μ L) were mixed with 4 mL of buffered dye
248 solution and incubated at 25 °C for 10 mins before absorbance was measured at 630 nm.
249 Linear standardisation of both methods was carried out using known concentrations of
250 bovine serum albumin and human serum albumin [Sigma- Aldrich, Poole, UK],
251 respectively. Serum globulin was estimated as the difference between total serum
252 protein and serum albumin; thereafter albumin and globulin ratio was subsequently
253 calculated.

254

255 Plasma glucose was measured through the oxidase-peroxidase method (Trinder, 1969).
256 Plasma samples (20 μ L) were incubated with 3 mL of oxidase-peroxidase stock solution
257 (16 mg 4-aminoantipyrine, glucose oxidase (1800 units, source: fungal, EC 1.1.3.4)
258 [Sigma-Aldrich, Poole, UK.], peroxidase (100 units, source: horseradish, EC 1.11.1.7)
259 [Sigma-Aldrich, Poole, UK.], 105 mg phenol, and 0.01% (v/v) Tween-20, made up to
260 100 mL phosphate buffer (100mM, pH 7.0) for 15 mins at 37 °C. This reaction was
261 ended by rapid cooling and absorbance (505 nm) was immediately measured. A linear
262 calibration curve was created using known concentrations of glucose standards [Sigma-
263 Aldrich, Poole, UK],

264

265 Plasma sodium and potassium measurements were carried out on a Corning 480 clinical
266 flame photometer [Corning, New York, USA]. Plasma samples (20 μ L) were diluted
267 fivefold using pure water (18.3 Ω M) before being analysed. Plasma triglyceride levels
268 were measured using a commercial colorimetric assay kit [Cayman chemicals Co., MI,
269 USA]. A commercial assay kit was used [Sigma-Aldrich, Poole, UK] to determine
270 plasma ammonia content.

271

272 Lipid peroxidation activity in the blood plasma was determined using the Gerard-
273 Monnier, Erdelmeier, Regnard, Moze-Henry, Yadan & Chaudiere (1998) 1-methyl-2-
274 phenylindole colorimetric method [Sigma-Aldrich, Poole, UK]. Briefly, 650 μ L reacting
275 agent (64 mg 10.3mM 1-methyl-2-phenylindole was dissolved in 30 mL acetonitrile,
276 mixed with 0.04408g 2,6-Di-*tert*-butyl-4-methylphenol dissolved in 10mL methanol)
277 was added to a 200 μ L plasma sample, together with 100 μ L 37% hydrochloric acid.
278 This was vortex mixed and incubated in a water bath at 45 $^{\circ}$ C for 40 mins. The reaction
279 was ended through immediate cooling in ice and centrifuged at 13,000 rpm for 10 mins.
280 The supernatant was spectrophotometrically read at 586 nm. A standard curve was
281 created through serial dilutions of diluted 1,1,3,3-tetraethoxypropane standard solution
282 (16.5 μ L of 10 mM 1,1,3,3-tetramethoxypropane in 20 mM Trizma-Base buffer pH 7.4,
283 added to 10 mL Trizma-HCL buffer).

284

285 **Statistical analysis**

286 Results were expressed as mean values and reported with pooled standard error (S.E.).
287 Datasets were analysed using nested (Treatment diet (replicate tank)) or one-way
288 analysis of variance (ANOVA) and significant differences among treatment means were
289 determined using *post-hoc* Tukey test. Statistical significance was only considered when

290 $P < 0.05$. All statistical analysis was performed on Statistical Package for the Social
291 Sciences-SPSS v16.0 [Manugistics Inc. Rockville, MD, USA].

292

293 **Results**

294 **Growth performance**

295 Fish fed on the ragworm inclusion (RW+FM) diet demonstrated a significantly superior
296 growth performance than those on all other diets ($P < 0.001$). This was shown by a
297 fivefold increase in weight gain, attainment of the lowest FCR, and the highest SGR and
298 PER (Table 3). Compared to the FM CTRL diet, the difference between the final mean
299 weight on RW+FM diet was 34% higher ($P < 0.001$). The silkworm pupae inclusion
300 (SWP+FM) diet *also* resulted in a significant increase in weight gain, producing a 12%
301 higher final mean weight ($P < 0.001$) when compared to the control diet. A combined
302 diet with both invertebrate meals (SWP+RW+FM) did not demonstrate any enhanced
303 morphometric parameters or growth performance indices when compared with the
304 control treatment (*Post-hoc Tukey*, $P > 0.05$).

305

306 **Basic haematological and immunological parameters**

307 Carp fed with either SWP+FM, RW+FM or SWP+RW +FM diet, produced higher
308 packed cell volume values than those on the control diet (Table 4). However, this was
309 found not to be significantly different ($P = 0.425$). Similarly, other basic haematological
310 parameters in the experimental carps were unaffected by the addition of invertebrate
311 meals into their diets ($P > 0.05$). Likewise, immunological parameters-lysozyme activity

312 and alternative complement pathway activity (ACH₅₀) showed no apparent significant
313 differences between the dietary groups. The exception was the SWP+FM dietary
314 treatment group, in which ACH₅₀ was significantly enhanced by approximately 41%
315 compared to the control diet (FM CTRL, $P=0.041$).

316

317 **Haematological biochemistry parameters**

318 Examination of total protein, albumin, globulin and albumin:globulin levels and plasma
319 Na⁺ and K⁺ concentrations showed no significant differences between treatment groups
320 ($P>0.05$, Table 5). Although, all experimental invertebrate meal diets were associated
321 with marked increases in fish glucose levels, there were no significant differences
322 compared to the control treatment ($P=0.333$). Blood ammonia levels showed no
323 statistically significant differences between the FM CTRL, SWP+FM and SWP+RW
324 +FM diets ($P>0.05$). However, experimental carp fed on RW+FM diet responded with
325 30% lower blood ammonia levels when compared to other dietary treatments ($P=0.009$).
326 Plasma triglyceride levels were significantly higher in fish that were fed either the
327 SWP+FM or RW+FM diets when compared to other experimental diets ($P=0.004$).
328 Levels of malondialdehyde in the blood were found to be highly variable between
329 sampled individuals (Table 5). Consequently, these differences were statistically
330 insignificant ($P=0.454$).

331

332 **Discussion**

333 The present study demonstrates that in comparison with a high BV protein in the form
334 of fishmeal, either ragworm or silkworm pupae meals can support good growth in

335 addition to enhancing growth performance in mirror carp under laboratory conditions.
336 Most notably inclusion of ragworm meal led to the highest growth performance
337 amongst the diet treatments. The results in the current investigation were favourably
338 comparable to those obtained by Salze *et al.* (2010) with juvenile Cobia (*Rachycentron*
339 *canadum*), who reported improvement in growth performance and feed efficiency when
340 fish were fed with 30% inclusion of ragworm diet. Rangacharyulu *et al.* (2003) reported
341 a 13% higher weight gain in *Catla catla*, mrigal (*Cirrhinus mrigala*) and rohu (*Labeo*
342 *rohita*) when 100% fishmeal was replaced by silkworm pupae silage. A previous study
343 in common carp (*Cyprinus carpio*) revealed it was possible to maintain growth
344 performance with a 50% silkworm pupae inclusion diet (Nandeesh, Srikanth,
345 Keshavanath, Varghese, Basavaraja, Das 1990). However, higher dietary inclusion
346 levels had negative impacts on juvenile Jian carp (*Cyprinus carpio*) physiology, such as
347 enhanced oxidative stress, occurrence of irregular-shaped intestinal structure, and
348 decreased protease and liver enzyme activities as reported recently by Ji *et al.* (2015).
349
350 Studies using other species of invertebrate meals have also resulted in improvements in
351 growth parameters measured for fish. These include the partial replacement of fishmeal
352 with mealworm (*Tenebrio molitor*) fed to African catfish (*Clarias gariepinus*) (Ng *et al.*
353 2001), and earthworm meal (*Eisenia foetida*) in rainbow trout (*Oncorhynchus mykiss*)
354 (Velasquez, Ibanez, Herrera & Oyarzun 1991). The rising costs associated with the
355 increasing demand for fishmeal could potentially result in the use of invertebrate meals
356 in fish feed production becoming economically viable (Sánchez-Muros *et al.* 2014).
357
358 An imbalance of the amino acid profile in fish diets can often result in decreased protein
359 retention efficiency and reduction of growth performance (Kaushik & Seiliez. 2010).

360 Moreover, excess supply of even a single amino acid can lead to enhanced amino acid
361 deamination and oxidation. This is in the case with glutamic acid, which is abundantly
362 found in wheat gluten based diets, resulting in a measurable increase liver glutamate
363 dehydrogenase activity and consequently raised ammonia production and ammonia
364 plasma concentration (Robaina, Corraze, Aguirre, Blanc, Melcion, & Kaushik, 1999;
365 Moyano, Cardenete & De La Higuera 1991). Furthermore, excessive supply of protein
366 relative to energy content can trigger protein catabolism to meet the energy demand
367 (McGoogan & Gatelin, 1999). Blood ammonia is the main excretory nitrogenous
368 product in fish, and in turn is excreted into the environment through the gills
369 (Weihrauch, Wilkie & Walsh 2009). The present study on carp resulted in a significant
370 reduction in ammonia production in fish fed the diet with the ragworm inclusion. From
371 this it can be inferred that there would be a reduction in ammonia concentrations in the
372 tank water, although these were not specifically measured. The implication is that
373 ragworm could be considered as a less polluting feed ingredient than fishmeal for mirror
374 carp under intensive rearing conditions and of significance to related ornamental fish
375 kept in tanks. The decrease in plasma ammonia and increase in protein efficiency ratio
376 suggest that the ragworm inclusion diet has an amino acid profile that is more
377 compatible with the dietary requirements of fast growing mirror carp than that provided
378 by the reference fishmeal-based diet, and thus better meets the profile for enhanced
379 protein accretion and consequently growth performance. The amino acid composition of
380 the diets (Table 2) indicated that a number of Essential Amino Acids, (EAA) such as
381 histidine, lysine, valine, methionine, and isoleucine were all lower for the reduced
382 fishmeal dietary treatments when substituted with both invertebrate based meals and a
383 blend. Fishmeal can present an excessive level of specific essential amino acids and
384 might not be as close to the 'ideal protein' concept for carp. For instance, in the

385 fishmeal control diet, lysine was determined as 3.01% and the NRC requirement for
386 common carp is stated as being 2.2% of the diet (5.7% of the Crude Protein CP, NRC,
387 2011). The invertebrate containing diets were closer in lysine (2.3-2.7%) to the
388 requirement of carp and may have been better assimilated. This may be true of other
389 amino acids and the overall requirement pattern compounded by the individual apparent
390 amino acid digestibility coefficients for each ingredient. These were not measured in the
391 present study due to the technical difficulties in acquiring faecal material from carp.
392 Longvah *et al.* (2011) used a rat model to assess protein quality and essential amino
393 acid score for silkworm meals and found that leucine scored low for this ingredient
394 compared to standard reference proteins such as hen egg and casein. However, leucine
395 did not appear to be rate limiting in the present diets for carp.

396

397 Both silkworm and ragworm meals have a higher lipid content than fishmeal. The lipid
398 content in the silkworm meal found in this study was similar to that previously reported
399 (Frye & Calvert, 1989; Rao, 1994; Hossain, Nahar, & Kamal 1997; Hertrampf &
400 Piedad-Pascual, 2000; Pereira, Ferrarese-Filho, Matsushita & De Souza. 2003).
401 Elevated lipid content in a feed component can make it difficult to formulate high
402 protein diets with a low to moderate lipid content, and would be a constraint for the low
403 oil diets required for carp production.

404

405 Triglyceride (TG) is one of the primary lipid storage compounds in fish and a principal
406 energy reserve in carnivorous species; it is also deemed important in overwintering carp
407 (Guillaume, Sadisivam, Bergot & Metailler, 2001). As well as being found within major
408 organs, such as the gastro-intestinal tract and liver, it is also abundantly located in the
409 blood lipid fraction, reflecting dietary status and history (Groff & Zinkl, 1999).

410 Although diets produced in this study were formulated to be *iso*-lipidic and *iso*-
411 energetic, haematological analysis revealed elevated blood plasma TG levels in mirror
412 carp fed on either of the single invertebrate meal inclusion diets. The increased growth
413 may explain the increased plasma TG levels in these fish, as the TG may have been
414 mobilised into the bloodstream via lipolysis from adipose tissue to be utilised to support
415 fish growth. Conversely, TG levels were not elevated in fish fed the mixed invertebrate
416 diet potentially because they did not display an associated enhanced growth. As dietary
417 lipid levels may be reflected in both whole body and fillet composition, it would be
418 prudent to carry out follow-up studies to examine whether the observed blood
419 triglyceride increase would significantly affect the fish fillet lipid composition of larger
420 carp reaching harvest size (Ferreira, De Araujo, Costa, Rosa, Figueiredo & Murgas
421 2011).

422
423 In many marine invertebrates, lipid composition is primarily comprised of the omega-3
424 series, eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) (Busarova &
425 Isay, 1984; Bischoff, Fink, & Wallera 2009), and in comparison, silkworm pupae are
426 rich in omega-3 α -linolenic acid (ALA) (Tomotake, Katagiri & Yamato 2010). The
427 combination of both high oil content and polyunsaturated fatty acid rich profile can lead
428 to increased susceptibility to lipid peroxidation in the invertebrate meal (Chen, Zhu,
429 Han, Yang, Lei, & Xie 2011). It can be assumed from the current study that the change
430 in lipid source did not in fact alter the level of lipid peroxidation as levels of the by-
431 product malondialdehyde, indicative of oxidative stress, remained unchanged. This
432 implies that lipid quality in both meals is relatively stable in terms of lipid oxidation
433 within these experimental diets, although the fatty acid profiles were not directly
434 measured in the present study.

435

436 The quality of lipids present in fish feeds can alter fillet quality, through physical and
437 organoleptic attributes (Ng & Bahurmiza, 2009). Besides lipids and their oxidation
438 products, other compounds present in feed ingredients can further modify quality. For
439 example, silkworms (*Bombyx mori*) are able to sequester flavonoids and terpenoids
440 compounds from their primary diet of mulberry (*Morus alba*) leaves (Zhou, Yang,
441 Chen, Lou, Zhang, Chen, Wang, Yu, Yu, Li & Zhong 2008). These compounds have
442 been linked with the strong odours associated with the pupae meal and can potentially
443 accumulate in fish muscle tissues. Moreover, a previous study has cited measureable
444 changes in fillet characteristics and in particular the organoleptic qualities leading to
445 'off flavour' in common carp fed experimental diets containing silkworm pupae meal
446 (Hora & Pillay, 1962). On the other hand, the presence of low molecular weight
447 bromophenol compounds in ragworms may serve as a functional feed component.
448 Found naturally in marine fish and shellfish, several of these compounds have been
449 shown to contribute to their distinct marine/ocean like flavour, as well as enhancing
450 existing flavours (Whitfield, Drew, Helidoniotis & Svoronos 1999; Ma, Chung, Ang &
451 Kim 2005). Further studies should be conducted in evaluating whether this novel meal
452 would have beneficial organoleptic properties for farmed fish. This would be
453 particularly relevant to freshwater fish such as carp that are typically grown in closed
454 systems (e.g. recirculated aquaculture systems). These systems are often rich in
455 cyanobacteria and actinomycetes, and metabolic by-products (e.g. geosmin) originating
456 from these organisms can accumulate in the fish muscle tissue causing a muddy flavour
457 (Robertson, Jauncey, Beveridge & Lawton, 2005). Through the use of ragworm meal in
458 the diet, it may be possible to mask this undesirable flavour by the presence of
459 bromophenols.

460

461 The current investigation with mirror carp showed enhanced alternative complement
462 pathway activity when fish were fed with the silkworm pupae inclusion diet. A possible
463 explanation for the improved immunity is the presence of long-chain polysaccharides in
464 the pupae meal. Like many invertebrates, the pupae of the silkworm have a cuticle
465 comprised mainly of chitin and chitosan, which gives rise to its structural integrity.
466 When the present study is compared to research conducted by Gopalakannan & Arul
467 (2006) and Lin, Pan, Luo & Luo (2011) with inclusion of extracted chitin and chitosan
468 in common carp (*Cyprinus carpio*) diets, an opposite trend was found, where lysozyme
469 activity was increased and alternative complement activity was decreased. However,
470 chitin was not directly measured in the present study, but would have been of interest to
471 help explain these effects.

472

473 **Conclusion**

474 Invertebrate meals may offer a sustainable feed material for farmed and ornamental
475 cyprinids such as carp, and furthermore, it can be sourced from waste-streams such as
476 silk production in Asia and polychaete culture in temperate zones. In the present
477 investigation, inclusion of either silkworm pupae or ragworm meal in diets for mirror
478 carp provides preliminary evidence that these novel raw materials can enhance growth
479 performance. In summary, this study has demonstrated that up to 7.5% of the dietary
480 protein content (24% of the fishmeal in experimental diets) can be replaced with either
481 terrestrial or marine derived invertebrate meals in order to sustain growth performance
482 of mirror carp (*Cypinus carpio*). The results also show there was no obvious detrimental
483 effect on selected blood parameters associated with health status. However, favourably

484 modulated components of the alternative complement pathway and decreased plasma
485 ammonia levels were indicative of reduced excretion of ammonia. The functional
486 effects of marine invertebrate meal on fish performance and health may advocate their
487 role as beneficial supplements to provide further 'added value' to the diet. Future work
488 should be directed towards characterising the effects of such novel protein sources for a
489 wider range of fish species within different production systems and phases of
490 development. Further, it may be possible to isolate the specific components and
491 metabolites in these novel ingredients responsible for their bioactive roles to optimise
492 utilisation in commercial diets for both aquaculture and the ornamental fish industry.

493 **Acknowledgments**

494 The research study was part funded by WALTHAM Centre for Pet Nutrition (Mars
495 Petcare Inc.). The authors would like to acknowledge Ben Eynon and Andrew Atfield
496 for their assistance during the feed trial and haematological analysis. Furthermore,
497 acknowledgement is also given to Liz Preston and Natalie Sweet for their technical
498 assistance in the nutritional analysis. The authors would to thank Dr Richard Fitzgerald
499 and Dr Majbritt Bolton-Warberg for their editorial assistance.

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Table 1: Proximate composition of the major feed components used in the experimental diets ($n=4$, %, dry matter).

Parameters	Herring Fishmeal (<i>Clupea harengus</i>)	Silkworm pupae meal (<i>Bombyx mori</i>)	Ragworm meal (<i>Nereis virens</i>)
Moisture	6.4	6.2	6.1
Crude protein	66.3	47.3	50.8
Crude Lipid	7.8	33.9	15.0
Crude Ash	13.0	3.1	7.9
Gross energy; MJ Kg ⁻¹	18.8	25.9	21.8

Table 2: Diet composition and proximate analysis of experimental diets (%).

Parameters	Diets			
	FM CTRL	SWP+FM	RW+FM	SWP+RW+FM
Diet composition				
Fish meal ^a	44.34	33.51	32.00	32.11
Silkworm pupae ^b	-	13.39	-	6.98
Ragworm meal ^c	-	-	14.80	8.00
Corn starch ^d	35.72	35.10	33.99	31.93
Lysamine [®] pea protein ^e	7.00	7.00	7.00	7.00
Glutalys [®] wheat gluten ^e	7.00	7.00	7.00	7.00
Mineral premix ^f	2.00	2.00	2.00	2.00
Sunflower oil ^g	2.00	2.00	2.00	2.00
Fish oil ^h	1.94	-	1.20	-
Calcium chloride ^d	-	-	-	2.98
Antioxidant ⁱ	0.50	0.50	0.50	0.50
Proximate composition¹				
Moisture	5.4	5.7	6.0	5.6
Crude protein	38.0	36.5	37.2	39.7
Crude Lipid	8.3	9.4	8.5	9.6
Crude Ash	7.3	6.5	6.8	7.4
Gross energy; MJ Kg ⁻¹	19.4	20.0	19.5	19.3
Essential Amino Acid composition				
Lysine	3.01	2.30	2.72	2.44
Methionine	1.11	0.85	0.99	0.89
Met + Cys	1.25	0.90	0.90	0.85
Arginine	2.74	2.56	2.43	2.48
Histidine	0.97	0.78	0.92	0.84
Threonine	1.68	1.56	1.60	1.58
Tryptophan	0.42	0.40	0.43	0.42
Leucine	3.46	3.41	3.22	3.32
Isoleucine	1.82	1.72	1.70	1.71
Phenylalanine	1.79	1.77	1.78	1.79
Valine	2.39	2.17	2.25	2.20

FM, Fishmeal (*Clupea harengus*); SWP, Silkworm pupae (*Bombyx mori*); RW, Ragworm (*Nereis virens*)

^a Scottish Fish meal 70, United fish products Ltd., UK.

^b Freeze dried silkworm pupae, Medikoi silkworm pupae, NT laboratories, UK.

^c Freeze dried ragworm meal, Seabait Ltd., UK.

^d Laboratory grade, Sigma –Aldrich Company Ltd., UK.

^e Roquette Frères, France.

^f Vitamin/Mineral premix, Premier Nutrition Products Ltd., UK. (Manufacturer analysis: Ca-12.09 %, Ash-78.71 %, Na-8.86 %, Vitamin A-1.0 µg kg⁻¹, Vitamin D3 0.10 %, Vitamin-E 7.0 g kg⁻¹, Cu-250 mg kg⁻¹, Mg 15.6 g kg⁻¹ and P 5.2 g kg⁻¹)

^g Sunflower oil, Costcutter Supermarkets Group Ltd., UK.

^h Epanoil, Sevenses Ltd., UK.

ⁱ Barox plus liquid (Antioxidant: Ethoxquine and BHT), Kemina Europa N.V., Belgium.

Table 3: Growth performance and feed utilisation of mirror carp (*C. carpio*), fed on different experimental diets (\pm S.E. $n= 3$).

Parameters	Diets				P-Value
	FM CTRL	SWP+FM	RW+FM	SWP+RW+FM	
Initial mean weight; g fish ⁻¹	14.93 \pm 0.19	14.63 \pm 0.09	15.07 \pm 0.13	14.90 \pm 0.05	0.132
Final mean weight; g fish ⁻¹	48.89 \pm 0.99 ^a	55.25 \pm 0.69 ^b	74.38 \pm 1.07 ^c	49.23 \pm 0.51 ^a	<0.001
Mean weight gain; g fish ⁻¹	33.96 \pm 0.77 ^a	40.62 \pm 0.72 ^b	60.83 \pm 2.57 ^c	34.34 \pm 0.48 ^a	<0.001
Specific Growth Rate, SGR; % day ⁻¹	1.78 \pm 0.04 ^a	1.99 \pm 0.02 ^b	2.36 \pm 0.04 ^c	1.81 \pm 0.01 ^a	<0.001
Feed Conversion Ratio, FCR; g g ⁻¹	1.92 \pm 0.02 ^a	1.68 \pm 0.02 ^b	1.36 \pm 0.2 ^c	1.83 \pm 0.01 ^a	<0.001
Protein Efficiency Ratio, PER	0.85 \pm 0.02 ^a	1.03 \pm 0.02 ^b	1.38 \pm 0.03 ^c	0.90 \pm 0.01 ^a	<0.001

Fish meal, FM; silkworm pupae (*Bombyx mori*), SWP; ragworm (*Nereis virens*), RW. Alternative complement activity, ACH50. Different superscript letters in the same row indicates significant difference ($P<0.05$).

Table 4: Basic haematological and immunological parameters of mirror carp (*C. carpio*), when fed different experimental diets (\pm S.E. $n= 3$).

Parameters	Diets				P-value
	FM CTRL	SWP+FM	RW+FM	SWP+RW+FM	
Packed cell volume; %	36.06 \pm 1.95	43.00 \pm 2.23	41.44 \pm 2.00	39.33 \pm 0.71	0.425
Haemoglobin; g dL ⁻¹	11.24 \pm 0.38	12.56 \pm 0.44	13.43 \pm 0.38	12.63 \pm 0.22	0.839
Erythrocytes; 10 ⁶ μ L ⁻¹	1.14 \pm 0.08	1.14 \pm 0.04	1.16 \pm 0.12	1.00 \pm 0.15	0.771
Lysozyme activity; units mL ⁻¹	28.44 \pm 3.97	28.44 \pm 2.70	27.56 \pm 3.43	32.89 \pm 4.26	0.888
ACH50; units mL ⁻¹	79.21 \pm 7.39 ^a	111.52 \pm 9.70 ^b	89.80 \pm 15.96 ^a	84.28 \pm 17.62 ^a	0.041

Fish meal, FM; silkworm pupae (*Bombyx mori*), SWP; ragworm (*Nereis virens*), RW. ACH50, alternative complement activity. Different superscript letters in the same row indicates significant difference ($P<0.05$).

Table 5: The biochemical haematology of mirror carp after being fed with experimental diets (\pm S.E., $n= 3$).

Parameters	Diets				P-value
	Control	SWP+FM	RW+FM	SWP+RW+FM	
Total Protein; g dL ⁻¹	1.36 \pm 0.02	1.39 \pm 0.03	1.51 \pm 0.03	1.37 \pm 0.02	0.161
Albumin; g dL ⁻¹	0.42 \pm 0.03	0.46 \pm 0.03	0.48 \pm 0.01	0.48 \pm 0.01	0.219
Globulin; g dL ⁻¹	0.97 \pm 0.04	0.87 \pm 0.04	0.92 \pm 0.06	0.89 \pm 0.03	0.529
Albumin/globulin ratio	2.50 \pm 0.34	2.02 \pm 0.27	1.92 \pm 0.15	1.86 \pm 0.10	0.232
Glucose; mg dL ⁻¹	79.86 \pm 1.75	93.31 \pm 2.97	93.50 \pm 3.33	90.41 \pm 1.93	0.333
Na ⁺ ; mmol L ⁻¹	116.39 \pm 0.25	125.00 \pm 1.71	118.98 \pm 1.47	116.94 \pm 0.72	0.901
K ⁺ ; mmol L ⁻¹	2.05 \pm 0.001	2.06 \pm 0.05	2.15 \pm 0.03	2.13 \pm 0.02	0.198
Ammonia; mg dL ⁻¹	21.37 \pm 2.24 ^a	20.60 \pm 1.41 ^a	14.56 \pm 1.21 ^b	24.37 \pm 2.01 ^a	0.004
Triglyceride; mg dL ⁻¹	75.37 \pm 7.10 ^a	99.67 \pm 13.21 ^b	102.64 \pm 17.81 ^b	64.62 \pm 3.72 ^a	0.009
Malondialdehyde; μ M mL ⁻¹	0.50 \pm 0.03	0.52 \pm 0.04	0.58 \pm 0.05	0.51 \pm 0.01	0.454

Fish meal, FM; silkworm pupae (*Bombyx mori*), SWP; ragworm (*Nereis virens*), RW.
Different superscript letters in the same row indicates significant difference ($P<0.05$).