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1 **The use of biorefinery by-products and natural detritus as feed sources for**

2 **Oysters (*Crassostrea gigas*) juveniles**

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6

7 **Abstract**

8 New research is currently underway to explore the potential of macroalgae for the production of
9 biofuels. Marine biofuels in general and macroalgae in particular, offer a number of advantages over
10 terrestrial biofuels including reduced competition for freshwater resources and for land use. Sugars
11 can be extracted from macroalgae and processed into biofuels by anaerobic digestion and
12 fermentation. This process generates significant waste biomass, which, if used, could improve the
13 economic sustainability of the biorefinery sector. Bivalves' aquaculture relies heavily on the
14 production of unicellular algae to feed juvenile individuals and this can represent a bottleneck for the
15 bivalve industry especially in locations where sunlight is limited. Previous research explored the use
16 of macroalgae derived digestate as alternative or integrative feed for juvenile bivalves, exploiting the
17 notion that organic particulate matter (detritus) is an integral part of this animal class natural diet. The
18 prospect of using waste products from the emerging biorefinery industry to solve a bottleneck for
19 aquaculture businesses and, by so doing, improving profitability of both, is an exciting one. In this
20 paper we describe the main nutritional profiles (Protein, Lipid, Carbohydrates and Fatty acids) of the
21 tested diets and investigate the potential for the use of a biorefinery a by-product as replacement
22 option for bivalves' production, by benchmarking it against aquaculture industry standards (live
23 microalgae and commercially available algae paste) and natural detritus constituted by farmed sea
24 urchin digesta. Both the digestate and the natural detritus supported the survival and growth of bivalve
25 spat, especially when used at 50% inclusion rate, over the course of 4-week preliminary trials. Data

26 suggest that a synergistic effect of the nutritional profiles of the diets employed may underpin the
27 observed results.

28 **Introduction**

29 Aquaculture is the fastest food production sector globally and the industry was worth US\$144.4
30 billion in 2012 (FAO, 2014). 90% of the industrial finfish and shellfish aquaculture producers have
31 juvenile or larval life stages that are micro-planktivorous (Duerr et al., 1998) and therefore would
32 greatly benefit from advances in early feeding protocols and products. Hatchery production of
33 bivalves is particularly reliant of constant and cost-effective production of unicellular algae.
34 Consequently, there is a pressing need in the production of bivalve juvenile to develop an inexpensive
35 and reliable feed that alleviates the reliance on live microalgae, a bottleneck of the bivalve industry
36 which constitutes as much as 30% of the overall spat production cost (Coutteau et al., 1994). In
37 addition to the financial aspect of producing microalgae, this process is also highly technical and
38 labour intensive, and the unpredictable growth of microalgae and the susceptibility of the culture to
39 contamination, has spurred interest in the development of more consistent and reliable alternative. At
40 present several species of live microalgae are utilised in the feeding of bivalve juveniles (Spolaore et
41 al., 2006). In traditional outdoor algae production systems it has proven difficult to maintain a
42 monoculture and successful growth is limited to regions with suitable temperature and sufficient
43 sunlight (Persoone, 1980). For these reasons more controlled and consistent systems that could be
44 utilised anywhere with a suitable power source were developed. Photobioreactors of various layouts
45 have been designed to produce highly controlled monocultures of algae for feed and for biofuels.
46 While these designs are often very successful at a laboratory scale it has generally been challenging to
47 scale them up to a commercial scale due to the relative decrease in illumination per unit area and
48 therefore an increased energy cost to adequately illuminate the microalgae (Ugwu et al., 2008). It
49 must also be noted that for the mass cultivation of algae, a large area is often needed and this
50 represents a common shortfall in many developed countries and has led scientists to investigate
51 alternative food sources for hatchery bivalves' production. An ideal replacement diet must be
52 nutritionally complete whilst being easily assimilated and absorbed. It must also exhibit
53 characteristics such as a long shelf life, an appropriate particle size for ingestion and a high retention
54 of its nutrients. Crucially, it must be less expensive to produce than current methods of microalgae

55 production. Currently, there are a number of alternatives to growing live microalgae available to the
56 aquaculture industry such as concentrated preparations of preserved non-viable microalgae
57 (PNVMA), yeasts and bacteria (Knauer and Southgate, 1999). However, these have met with limited
58 success either due to cost of production, their physical properties or their nutritional content.
59 Therefore, the development of a diet to replace unicellular algae has a significant industrial value
60 (Schiener et al., 2015). The role of macrophyte detritus as a food source in many ecosystems is well
61 documented [Charles, 1993; Nagelkerken et al., 2008]. It has been long established that bivalves
62 readily absorb Kelp detritus and its associated bacteria, which suggests that it can be an important
63 food source for this animal class (Stuart et al., 1982). In previous studies successes have been
64 observed when utilising a single cell detritus (SCD) feed produced from the degradation of marine
65 macrophytes, (Uchida, 1996; Uchida and Murata, 2002; Perez Camacho et al., 2004). Degradation of
66 macroalgae can be achieved through a multitude of processes involving exposure to a combination of
67 proteolytic, alginolytic and cellulolytic enzymes, pH manipulation and bacteria. The size of the
68 particles available after degradation and processing is below 20µm, which is analogous with typical
69 dietary phytoplankton species and suggests its usefulness as a nursery feed for molluscs. Early
70 studies (Uchida et al., 1997a; Uchida et al., 1997b) confirmed this and found that SCD from thalli of
71 *L. japonica* degraded using the marine bacteria was a viable food source for *Artemia salina* nauplii
72 and, more recently, SCD from *Porphyra haitanensis* was found to be a successful substitution diet for
73 nursery production of the tropical oyster *Crassostrea belcheri* (Tanyaros and Chuseingjaw, 2014).

74 Although the use of farmed macroalgae for biofuel production and the potential for modifying their
75 biochemical profile via environmental manipulation dates back to the 1980s (Rythers et al., 1981;
76 Bird and Benson, 1987), recently, the concept has seen an increased interest (Hughes et al., 2012;
77 Kraan, 2013) and it has been significantly developed to improve its economic viability. One further
78 significant improvement in the economic performance of biorefinery could be represented by the use
79 of the process's by-products as valuable feed sources for livestock, including marine bivalves. At
80 laboratory scale, the use of biorefinery by-product has been shown to have potential as bivalve feed,
81 mostly due to the feeding habit of this animal class, which includes particulate organic matter (POM)

82 as a significant component of its natural diet (Mann, 1988; Duggins et al., 1989). Therefore, these
83 digestates, or Single Cell Detritus (SCD), from marine macroalgae, obtained via enzymatic digestion,
84 have the potential to mimic the physical properties and biochemical profiles of natural particulate
85 organic matter and consequently fulfil, at least partially, bivalves' nutritional requirements. Indeed,
86 the elemental composition of macroalgae degraded via enzymatic saccharification and their potential
87 as a replacement for commercially available PNVMA has been recently described (Schiener et al.,
88 2015). With this study, we take this concept further and compare the biochemical composition and
89 suitability as oyster feed of the SCD produced by *S. latissima* enzymatic saccharification used in a
90 previous study (Schiener et al., 2015) with live microalgae as well as commercial algae paste.
91 Importantly, a comparison between biochemical composition and suitability as aquaculture feed
92 between biorefinery by-products and natural detritus is, to our knowledge, still lacking. The reduction
93 of macrophytes to a SCD product through acidic, bacterial, enzymatic and mechanical action can, in
94 fact, be associated to the animal digestive process. It could therefore be hypothesised that the
95 digestive action of a marine grazing herbivore would produce a product of similar composition to that
96 of "artificially" produced detritus. Sea urchins are one of the major consumers of macro-phytobenthos
97 and, as such, possess the potential to significantly contribute to the particulate organic matter fraction
98 in several marine ecosystems, providing an important link in the nutrients fluxes between the benthic
99 and pelagic domains. This study, therefore, assesses the viability of SCD produced via the digestive
100 action of the sea urchin *Paracentrotus lividus* compared to SCD produced by enzymatic
101 saccharification in an anaerobic digester, for the hatchery production of *Crassostrea gigas* spat, by
102 benchmarking these two novel diets against industry standards: live microalgae and commercially
103 available algae paste. This paper describes the growth, survival and biochemical composition
104 (Carbohydrates and Lipids) of juvenile oysters (*C. gigas*) and reports on the biochemical composition
105 of the tested diets (Proteins, Carbohydrates, Lipids and Fatty acids).

106

107 **Materials and Methods**

108 *General methods*

109 In this study six diets were trialled in triplicate; a live microalgae diet consisting of a 70:30% by algae
110 cell volume mix of *T. suecica* and *I. galbana* (MA), an algal paste diet (AP) supplied by Reed
111 Mariculture Inc. (Shellfish Diet 1800®), Single Cell Detritus produced by enzymatic saccharification
112 (SCD); natural detritus produced from *Paracentrotus lividus* faeces (UF); 50% MA-SCD and a 50%
113 MA-UF. The oysters were kept in 3 litre glass bowls in a static system with an air stone in each bowl
114 to maintain circulation and prevent settling of feed particles. Into each bowl was placed 700mg of spat
115 (approximately 150 individuals; wet weight 4.6 ± 0.2 mg; shell length 1.96 ± 0.44 mm) on a raised mesh
116 platform, to allow full circulation of water and feed to each individual. Water temperature of the
117 bowls was maintained at 16.3°C (+/- 0.8 SD) using manipulation of the ambient room temperature.
118 Where required feeds were converted into a liquid form by adding the dry feeds to either ambient
119 seawater or to the respective algae mix, algae paste was diluted with ambient filtered seawater as per
120 supplier instructions. The feed rations were administered in a pulse format of 24 separate feeds of
121 10ml once every hour. Daily Feed rations for each treatment were calculated and replenished once per
122 day. The Jebao DP 4 peristaltic pump was used to apply the hourly rations for each replicate. Bowls
123 were cleaned using warm fresh water and complete water change was conducted every three days.
124 Treatments were kept in a temperature controlled room and maintained on a photoperiod of 8 hours of
125 daylight and 16 hours of darkness. Rations of the live algae mix used to feed the MA, MA-SCD and
126 MA-UF treatments was calculated daily according to published methods (FAO, 2004).

127 Rations of the Shellfish Diet 1800® for the algae paste treatments were calculated based on the
128 manufacturer guidelines. Rations of both the SCD and UF diets were calculated based on a 40% of
129 oyster live weight per week in diet dry weight, in a way that the ration for these diets matched the
130 ration of both live microalgae and algae paste (FAO, 2004). Randomly picked 80 individuals from
131 each of the replicates were weighed to determine individual wet weight and were measured using
132 callipers to determine shell length. A mortality count was also undertaken on the same amount of
133 individuals per replicate. Oysters were considered to be dead when presenting open shells or showed
134 no dark coloration or mantle movement when observed under dissecting microscope.

135

136 *Preparation of the Diets*

137 The UF feed was produced from the faeces of *Paracentrotus lividus* fed to satiation with *S. latissima*
138 fronds. The faeces were collected soon after production in an effort to minimize nutrient leeching. The
139 wet faeces was sieved through a 200µm mesh to remove large uneaten particles and broken urchin
140 spines, it was then allowed to settle in tall 1 litre measuring cylinder and the supernatant was siphoned
141 off. The faeces were transferred to a shallow tray and allowed to air dry at room temperature (21°C),
142 any remaining spine fragments were removed during this process by hand while the faeces was still
143 moist. As soon as the faeces had dried sufficiently to be scraped from the tray as a paste it was freeze
144 dried to remove moisture. The dried faeces were then ground to a fine powder using a pestle and
145 mortar and stored in a desiccator. Using a fume-hood to minimize dust inhalation the fine powder was
146 sieved using a 20µm test sieve to ensure all particles were below 20µm and could be ingested by the
147 spat.

148 Live algae diet was a 70:30 mix of *Tetraselmis suecica* and *Isochrysis galbana* grown in sterile 20
149 litre carboys with the addition of f/2 medium. Algae Paste used was the Shellfish Diet 1800®
150 purchased from Reed Mariculture Inc. four days prior to the start of the trial.

151 To produce the SCD diet fronds of *Saccharina latissima* were treated using cellulosic and
152 hemocellulosic enzyme blends provided by Novozymes, Denmark (Schiener et al., 2015).
153 Approximately 13.00 ± 0.002 g of dried seaweed was added to 250ml Duran glass bottles with 100ml
154 of deionised water. The pH of the solution was adjusted to 5.2 with 10% HCl and the bottles
155 autoclaved at 121°C for 15 minutes. Once cooled to 45°C in a water bath, enzymes were added at 10%
156 NS 22086 (w w-1) and 1.2% NS 22119 (w w-1). Bottles were placed in an orbital shaker (New
157 Brunswick Scientific, Innova 4230) at 200 rpm and incubated at 45°C for 2 days. Following this, the
158 digested seaweed was centrifuged for 10 minutes at 3.200g and residue was washed with equal
159 volumes of deionised water before re-centrifugation. Washed solids were frozen at -20°C and vacuum
160 freeze dried to remove all moisture. The dried matter was then mechanically ground using a pestle and
161 mortar to reduce particles size and sieved through a 20µm mesh.

162

163 *Biochemical Analysis*

164 Each of the six diets was processed into a dry powder by centrifugation at approximately 5000rpm for
165 10 minutes, supernatant was drained and the remaining pellet was freeze dried and ground into a fine
166 powder. The MA-SCD and MA-UF dried diets were made by combining the respective dried powders
167 at a 1:1 ratio based on weight.

168 The lipid fraction of diets and oysters was extracted using procedures described by Folch (Folch et al.,
169 1957). In brief, samples were homogenized in the chloroform/methanol using a tissue disrupter (Ultra
170 Turax™, IKA Werke GmbH & Co. KG, Staufen, Germany), and 1 ml 0.88% KCl was added and the
171 homogenates mixed before centrifugation at 600 g for 5 min (Jouan C412, Pegasus Scientific Inc.,
172 Rockville, USA). The upper aqueous phase was aspirated and the solvent evaporated under a stream
173 of oxygen-free nitrogen (OFN). Lipid content was determined gravimetrically after desiccation
174 overnight. The total lipid extracts were re-dissolved at a concentration of 10 mg/ml in
175 chloroform/methanol (2:1, v/v) plus BHT. Fatty acid compositions of total lipid were determined by
176 gas chromatography according to standard protocols (Christie, 2003). Fatty acid methyl esters
177 (FAME) were prepared from total lipid by acid-catalyzed transesterification at 50 °C for 16 h with
178 extraction and purification by thinlayer chromatography as described previously (Ackman, 1980). The
179 FAME were separated and quantified by gas-liquid chromatography using a GC 8000™ series EL
180 980 GLC (Fisons instruments) equipped with a 30 m× 0.32 mm i.d., 0.25 µm capillary column (CP
181 Wax 52CB, Chrompak, London, U.K.) and on-column injection. Hydrogen was used as carrier gas
182 and temperature programming was from 50 to 150 °C at 40 °C min⁻¹ and then to 230 °C at 2.0 °C min⁻¹.
183 Individual methyl esters were identified by comparison with known standards and by reference to
184 published data (Ackman, 1980; Tocher and Harvie, 1988). Data were collected and processed using
185 Chromcard for Windows (version 1.19), and FAME quantified through a comparison with a
186 heptadecanoic acid (17:0) internal standard.

187 Carbohydrate content of the diets was measured using a Uvikon™ 860 spectrophotometer and
188 compared to a calibration curve generated from known quantities of glucose standard. Between 2.7
189 and 14.5 mg of whole, freeze dried oyster was used in each replicate. Between 2.8 and 8.4 mg of
190 dried, powdered diet was used in each replicate. Each treatment was analysed in triplicate. Solutions
191 made up of 2.5 ml deionised water, 1 ml of 5% phenol solution and 8 ml of concentrated sulphuric
192 acid in the necessary order and at the necessary time in the procedure. The absorbance of each
193 solution was read at 520 nm against a blank standard. From the calibration curve the mg of glucose
194 for each replicate can be determined and converted into total carbohydrate using the following
195 formula:

$$196 \quad \% \text{ total carbohydrate} = (\text{mg of glucose in sample} / \text{sample weight (g)}) \times 100$$

197 Protein of the diets was measured using the Kjeldahl analysis on a Tecator Kjeltec according to Lynch
198 and Barbano (1999). Between 71.9 and 276.9 mg of dried, powdered diet was used for each replicate,
199 all samples were analysed in duplicate. Two copper Kjeltabs and 5 ml of concentrated sulphuric acid
200 was added to each replicate before placing the tubes into a digestion block at 420°C for 1 hour. 20 ml
201 of deionised water was then added before allowing the mixed solution to distil using a Kjeltec™ 2300
202 analyser (FOSS).

203

204 *Statistical Analysis*

205 All analyses were carried out using the statistical package of Mini-tab 15.0 (Minitab Ltd., UK).
206 Normality and homogeneity of variance were ere confirmed using Kolmogorov–Smirnov test and
207 improved where necessary by either log or reciprocal transformations. Differences were tested using
208 one-way ANOVA, followed by the Tukey’s multiple comparison test to assess where significant
209 differences occurred. The non-parametric multivariate analysis ANOSIM (analysis of similarities)
210 was used to identify significant differences in the diets fatty acids profiles. SIMPER (similarity
211 percentage) test was used to identify which FAs were primarily responsible for the observed

212 differences (Carboni et al., 2013). Data were untransformed and Euclidian distance was used as the
213 metric. In all cases, significant differences were determined at $p<0.05$.

214

215 **Results**

216 *Oyster growth, survival and nutritional reserves*

217 The feeding trial showed that the oysters in all treatments have significantly grown during the trial
218 period ($p<0.05$) and that survival was generally high with no difference across treatments (Tab. 1).
219 However, the biofuel residue (SCD), the detritus produced from sea urchin faeces (UF) and
220 commercial algae paste (AP) were only marginally capable of supporting oyster spat growth when fed
221 on their own. Conversely, when both SCD and UF were used as 50% live algae substitute, significant
222 faster growth was observed (Fig. 1). This confirms the nutritional value of these residues as potential
223 bivalve diet supplement or partial replacement but not as standalone diets. Indeed, by the end of the
224 four weeks feeding trial, oysters fed the MA-UF diet had a significantly higher mean individual
225 weight compared to all other diets, including live microalgae (Fig. 1), suggesting that the nutritional
226 profile and/or the digestibility of the UF supplement should be further investigated as it appears to
227 provide a growth advantage. Although shell length at the end of the trial was significantly higher than
228 at the beginning, no significant differences were observed between the treatments (Tab. 1).

229 Oysters' carbohydrate and lipid content at the end of the trial period is given in Table 1. Data show
230 that individuals in every treatment accumulated nutrients reserves during the trial period, suggesting
231 that efficient feeding was achieved with the employed experimental system. No difference in lipids
232 and carbohydrates content were observed between the oysters fed the detritus based diets and
233 commercial algae paste. However, oysters fed MA had a significantly higher nutritional content
234 ($p<0.001$), indicating the higher long-term suitability of this diets as oyster feed.

235

236 *Biochemical composition of the diets*

237 Table 2 shows the protein, carbohydrate, lipid and fatty acids content of all tested diets. Significant
238 difference between protein content of the diets was observed ($p < 0.001$). The protein content of the
239 Single Cell Detritus produced by enzymatic saccharification (SCD), was significantly higher
240 ($30.45 \pm 0.40\%$) than any other diet. The second highest protein content was measured in the
241 commercial algae paste (AP) diet ($21.80 \pm 0.14\%$) and in the MA-SCD diet ($19.15 \pm 0.60\%$), whilst no
242 significant difference were observed between the remaining three diets. Ideal dietary protein content
243 for juvenile bivalves has been estimated to be between 13% for *R. decussatus* (Albentosa et al., 1996)
244 and 20% for *C. virginica* (Flaak and Epifano, 1978), although we can assume the requirement for *C.*
245 *gigas* is closer to the latter. The protein content of the two best performing diets, MA-UF and MA
246 had, however, the two lowest protein content of any diet.

247 The carbohydrate content of the MA-UF diet ($10.39 \pm 0.44\%$) and the AP diet ($10.788 \pm 0.94\%$) were
248 not significantly different. The remaining diets showed significant differences ($p < 0.05$). More
249 specifically, the detritus diets and their relative 50% mix with live microalgae had the highest
250 carbohydrates content compared to commercial algae paste and live microalgae. In particular, the
251 detritus produced from anaerobic digestion (SCD) contained almost 8 times the amount of
252 carbohydrates than MA.

253 Lipid content of the MA, SCD the MA-SCD diet and MA-UF diets did not differ significantly. Lipid
254 content of the AP diet, instead, was significantly higher than all the other diets ($p < 0.001$). The total
255 effect of lipid content of a diet on the growth of *C. gigas* spat has been found to be relatively
256 insignificant (Langdon & Waldock, 1981). This is consistent with the results presented here as the
257 higher lipid content of the AP diet was not matched by animal growth performances. Fatty acids
258 profiles of all the tested diets are presented in the non-Metric Multidimensional Scaling plot (Fig. 2).
259 From this one-way Anosim analysis of the dietary fatty acid profiles, it is clear that the detrital diets
260 (SCD and UF) presented a very distinct profile from the live microalgae and algae paste diet. (MA
261 and AP) Interestingly, however, when the former were mixed with live microalgae their fatty acid
262 profile was tightly clustering with the MA diet. The simpler analysis showed that the main fatty acid
263 responsible for the observed difference between MA and AP was 16:1n-7, which on its own

264 contributed for over 20% of the profiles differences, whilst n-3 and n-6 fatty acids only minimally
265 contributed to the difference. On the contrary the main fatty acids contributing to the differences
266 between detrital diets and AP and MA were of the n-3 group, mainly EPA and DHA.

267 Significant differences between diets were observed in the main fatty acids groups: saturated,
268 monounsaturated, n-6 polyunsaturated and n-3 polyunsaturated (Fig. 3). Saturated fatty acids were
269 observed to be in significantly higher amount in the UF diet ($49.41 \pm 0.77\%$) than all other treatments
270 ($p < 0.001$). Monounsaturated fatty acid (MUFA) content was highest ($p < 0.001$) in the SCD treatment
271 ($49.659 \pm 0.32\%$), but there was no significant difference between the MA-SCD diet ($34.48 \pm 1.66\%$)
272 and the UF diet ($35.758 \pm 0.32\%$). There was also no significant difference between the MA/UF and
273 the AP diet. The n-6 PUFAs content was significantly different between the diets ($p < 0.01$) and AP
274 and SCD showed the highest amounts. Finally, 3-n PUFAs were significantly higher in the MA diet
275 compared to all others ($p < 0.001$). Eicosapentaenoic acid (20:5n-3, EPA), Docosahexaenoic acid
276 (22:6n-3, DHA) and Arachidonic acid (ARA, 20:4n-6) and their respective ratios are considered
277 particularly important in animal physiology and, in many marine species, are considered to be
278 essential fatty acids (EFAs) that need to be provided by the diet (Knauer and Southgate, 1999; Tocher,
279 2003). Figure 4 shows the relative abundance of these important compounds in the tested diets. The
280 EPA content was significantly different across each diet ($p < 0.01$) with the AP diet showing a
281 considerably higher content than any other tested diet. The MA diet had significantly higher levels of
282 DHA compared to the other diets ($p < 0.01$). The UF diet contained a comparatively small amount
283 DHA, while the SCD diet did not contain any (Fig. 4). Juvenile *Cerastoderma edule* growth did not
284 change when fed a diet containing high levels of EPA and DHA when ARA was instead deficient; this
285 indicates that EPA and DHA may be the most crucial EFAs for juvenile bivalve growth (Reis Batista
286 et al., 2014). Importantly, bivalves do possess some ability to elongate and desaturate precursor fatty
287 acids such as 18:3n-3 into EPA and DHA, if only at low levels (Da Costa et al., 2015). This in turn
288 indicates that high levels of EPA and DHA may not be as important in marine bivalves as they are in
289 marine carnivorous fish.

290

291 **Discussion and Conclusion**

292 Both the digestate and the natural detritus supported the survival and growth of bivalve spat,
293 especially when used at 50% inclusion rate, over the course of this 4-weeks preliminary trial. Despite
294 these promising results, however, it is important to notice that the growth rate achieved by the
295 juvenile oysters fed MA-UF was only half of that commonly observed under commercial conditions
296 (*pers. obs.*) using commercial upwelling systems. This, in combination with the oysters' nutritional
297 reserves, strongly indicates that further research into these new potential feed replacements should be
298 conducted using commercial protocols before these results could up-taken by the industry. This is
299 particularly important considering that the use of static tanks with a low volume (2-4l) can lead to an
300 increased growth of bacteria which can contribute to the nutrition of the animals (Laing, 1987). The
301 effect of bacterial proliferation is not yet clear. In some circumstances the bacteria caused clumping
302 which inhibited ingestion (Langdon, 1983). However, clumping effect has also been found to serve as
303 an undefined food source with bacteria contributing significantly to the metabolic nitrogen
304 requirement of *C. virginica* in closed systems (Langdon and Newell, 1990).

305 The protein content of the two best performing diets, MA-UF and MA has shown the two lowest
306 content of any diet. This seems, therefore, to suggest that a protein content of approximately 9% was
307 sufficient under the trial conditions employed here. Nonetheless, full aminoacid profile would have
308 provided more clarity for the interpretation of these results. It is also worth noting that the interaction
309 of protein with other nutritional elements and the amino acid profiles of the diets was not analysed in
310 this study and may have been an important factor (Utting, 1986).

311 The biochemical analysis showed that the detritus produced from anaerobic digestion (SCD) and the
312 natural detritus (UF) contained almost 8 times the amount of carbohydrates than MA. Carbohydrate is
313 mainly utilised as an energy source by juvenile bivalves and acts to balance the utilization of protein
314 and lipid for biosynthesis and growth against catabolism for energy (Whyte et al., 1989). It has been
315 found that ingestion of carbohydrate is closely correlated with growth in *C. gigas* spat (Brown et al.,
316 1998), however this is not consistent with the results from this trial as the SCD diet contained

317 significantly higher amounts of carbohydrates than other diets although it wasn't the best performing
318 diet. This suggests that requirements may be fulfilled at lower levels, and that other nutritional factors
319 must be met to facilitate all potential growth. It is also possible that the detrital component of the MA-
320 SCD diet was not as palatable or digestible as the MA-UF diet and was therefore not ingested or
321 digested at the same rate. Furthermore, the increased carbohydrates content combined with a richer n-
322 3 fatty acid profile of the MA-UF diet could be at the root of the better growth performances of the
323 oyster fed this diet.

324 As expected, the three treatments that included the live Microalgae mix performed the best overall.
325 The MA and AP diets were intended to establish an industry consistent benchmark and it was not
326 anticipated that any diet would perform better than the live microalgae diet. Surprisingly, individual
327 wet weight of oysters fed the MA-UF was instead significantly higher than that of animals fed live
328 microalgae alone. This diet also outperformed both the SCD and the algae paste diets that were
329 previously shown to possess potential as live microalgae replacement in the hatchery production of
330 oyster juveniles (Schiener et al., 2015). These findings suggest that the MA-UF diet was either the
331 most nutritionally complete (i.e. more suitable carbohydrate content and fatty acid profile) and/or
332 most bioavailable. The AP diet showed similar levels of nutrients to the MA diet; however, growth in
333 the AP treatments was significantly slower. Likewise, the MA-SCD and MA-UF diets had very
334 similar nutritional profiles despite the MA-UF diet performing significantly better overall. This
335 suggests that beside nutrient density there is a much more complex range of parameters, such as
336 settling rate, ingestion rate and assimilation rate, that contribute to the success of a diet and highlights
337 the need for successive studies to ascertain the key factors that allowed the UF feed to be so
338 successful when used in conjunction with a multi-specific algal diet.

339 New research is currently underway to explore the potential of macroalgae for the production of
340 biofuels (Suutari et al., 2015) as hexose sugars can be extracted from macroalgae and processed into
341 biofuels by anaerobic digestion and fermentation (Goh and Lee, 2010; Chen et al., 2015). This process
342 generates significant waste biomass, which can, in theory, be utilised and further processed into an
343 SCD product. Sea urchin digestion process is still under-researched and the findings from this study

344 suggest that digestive enzymes and/or the microbiota associated with echinoderms digestive processes
345 could provide valuable information for the advancement on marine biomass exploitation and, at the
346 same time, produce residuals that may prove to be advantageous for the aquaculture industry.
347 Nonetheless, the actual economic implications of this hypothetical partnership are, difficult to
348 speculate due to both industries infancy and collaborative interdisciplinary research should be
349 conducted to evaluate the technical and economic scope of such initiative.

350

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Tables

Table 1.

Nutritional reserves, size (Shell Length) and survival of the oysters spat at the end of the 4 weeks experimental period (mean \pm sd; n=3). Superscripts indicate statistically significant differences.

Table 2.

Biochemical composition of the six tested diets (mean \pm sd; n=6). Superscripts indicate statistically significant differences.

Table 1

	Initial	MA	AP	SCD	UF	MA-SCD	MA-UF
Lipid (% tissue weight)	0.53±0.28 ^c	1.49±0.32 ^a	0.41±0.12 ^c	0.38±0.08 ^c	0.52±0.12 ^c	1.19±0.33 ^b	0.68±0.19 ^c
Carbohydrates (% tissue weight)	0.71±0.17 ^c	2.66±0.60 ^a	1.59±0.32 ^b	1.65±0.32 ^b	1.64±0.39 ^b	1.90±0.53 ^b	1.83±0.57 ^b
Shell Length (mm)	1.96±0.44 ^b	3.80±0.36 ^a	3.05±0.72 ^a	3.00±0.53 ^a	2.93±0.65 ^a	3.51±0.28 ^a	4.16±0.46 ^a
Survival (%)		97.7±3.13	95.36±5.15	95.85±4.80	93.98±6.16	93.91±5.22	94.83±4.80

Table 2

Diets	MA	AP	SCD	UF	MA-SCD	MA-UF
Proteins (% of dw)	8.11±0.73 ^d	21.80±0.14 ^b	30.45±0.40 ^a	9.33±0.06 ^c	19.15±0.60 ^b	8.47±0.29 ^d
Carbohydrates (% dw)	5.63±0.85 ^d	9.90±1.19 ^c	39.34±2.60 ^a	16.77±0.51 ^d	22.63±2.64 ^b	10.26±0.72 ^c
Carbohydrates/Protein	0.70±0.12	0.46±0.06	1.26±0.10	1.79±0.08	1.19±0.16	1.23±0.08
Lipids (% of dw)	5.48±0.71 ^b	12.56±0.16 ^a	6.07±0.40 ^b	3.46±0.84 ^b	5.63±0.49 ^b	4.02±1.10 ^b
Fatty Acids (% of total lipids)						
14:0	10.96±0.76 ^a	8.98±0.27 ^b	4.80±0.08 ^d	7.63±0.11 ^c	8.22±1.01 ^{bc}	10.40±0.86 ^a
iso 15:0	0.21±0.03 ^e	0.38±0.02 ^d	1.61±0.03 ^b	2.70±0.05 ^a	0.93±0.06 ^c	0.93±0.16 ^c
15:0	0.24±0.01 ^e	0.76±0.02 ^b	0.72±0.01 ^c	1.81±0.16 ^a	0.49±0.03 ^d	0.63±0.10 ^c
16:0	11.55±0.88 ^f	13.49±0.74 ^e	20.24±0.24 ^b	34.48±0.65 ^a	15.67±0.32 ^d	18.19±0.45 ^c
18:0	0.32±0.14 ^e	0.46±0.04 ^e	2.94±0.04 ^a	2.32±0.19 ^b	1.57±0.12 ^c	0.97±0.06 ^d
20:0	nd	nd	0.51±0.02 ^b	0.65±0.01 ^a	nd	0.20±0.02 ^c
Total saturated	23.29±1.65 ^d	24.31±0.64 ^d	31.21±0.33 ^b	49.81±0.65 ^a	27.39±0.95 ^c	31.41±0.88 ^b
16:1n-9+DMA	5.11±0.18 ^c	6.57±0.53 ^b	8.81±0.22 ^a	6.57±0.19 ^b	7.10±0.25 ^b	5.41±0.31 ^c
16:1n-7	1.86±0.10 ^f	12.53±0.29 ^a	11.19±0.22 ^b	9.79±0.22 ^c	6.35±0.43 ^d	4.27±0.17 ^e
18:1n-9	10.85±1.50 ^c	5.67±0.31 ^d	26.66±0.48 ^a	12.93±0.18 ^c	18.32±0.56 ^b	11.51±1.45 ^c
18:1n-7	2.69±0.11 ^c	0.95±0.05 ^e	2.36±0.05 ^d	4.92±0.13 ^a	2.47±0.05 ^d	3.41±0.18 ^b
20:1n-9	0.78±0.09 ^a	0.18±0.01 ^d	0.21±0.14 ^d	0.26±0.11 ^d	0.43±0.03 ^c	0.55±0.02 ^b
Total monounsaturated	21.68±1.28 ^d	26.10±1.13 ^c	49.31±0.45 ^a	34.85±1.10 ^b	34.84±1.06 ^b	25.41±1.04 ^c
18:2n-6	3.17±0.18 ^d	4.39±0.03 ^c	6.50±0.13 ^a	2.58±0.24 ^d	4.84±0.14 ^b	3.02±0.17 ^d
18:3n-6	0.09±0.01 ^d	1.09±0.04 ^a	0.34±0.01 ^b	0.11±0.01 ^d	0.22±0.01 ^c	0.09±0.00 ^d
20:4n-6	0.12±0.01 ^d	0.60±0.03 ^c	1.69±0.05 ^a	1.65±0.07 ^a	1.01±0.08 ^b	0.61±0.06 ^c
22:5n-6	1.52±0.05 ^b	2.22±0.14 ^a	nd	nd	0.78±0.09 ^d	1.10±0.05 ^c
Total n-6 PUFA	5.36±0.10 ^d	8.43±0.19 ^b	8.67±0.14 ^a	4.83±0.18 ^e	7.16±0.19 ^c	6.16±1.81 ^d
18:3n-3	12.26±1.43 ^a	4.19±0.14 ^d	2.55±0.07 ^e	2.53±0.05 ^e	7.43±0.61 ^c	9.44±1.16 ^b
18:4n-3	11.32±0.25 ^a	8.90±0.87 ^b	1.64±0.06 ^d	1.05±0.03 ^e	6.60±0.42 ^c	8.38±0.22 ^b
18:5n-3	5.06±0.41 ^a	0.74±0.09 ^d	nd	0.32±0.04 ^e	2.67±0.17 ^c	3.82±0.24 ^b
20:5n-3	3.12±0.10 ^b	13.84±1.29 ^a	1.20±0.03 ^f	1.72±0.14 ^e	2.19±0.09 ^d	2.66±0.013 ^c
22:6n-3	8.16±0.33 ^a	5.50±0.63 ^b	nd	0.21±0.02 ^d	4.29±0.33 ^c	6.07±0.25 ^b
Total n-3 PUFA	40.62±2.36 ^a	33.63±3.04 ^b	5.54±0.15 ^d	5.97±0.28 ^d	23.59±1.24 ^c	29.92±2.43 ^b
16;2	0.82±0.15 ^b	2.20±0.08 ^a	0.23±0.01 ^d	0.17±0.01 ^e	0.55±0.05 ^c	0.65±0.10 ^{bc}
16;3	0.33±0.43 ^c	1.55±1.48 ^a	1.16±0.04 ^b	0.47±0.04 ^d	0.69±0.05 ^c	0.18±0.02 ^e
16;4	5.90±0.21 ^a	0.81±0.15 ^d	nd	nd	2.76±0.26 ^c	3.94±0.19 ^b
15:0 DMA	0.53±0.03 ^e	0.79±0.07 ^d	1.04±0.11 ^b	1.89±0.65 ^a	0.84±0.04 ^c	0.76±0.23 ^{cd}
16:0 DMA	1.47±0.08 ^c	2.18±0.15 ^b	2.84±0.05 ^a	2.03±0.03 ^b	2.18±0.14 ^b	1.58±0.13 ^c
Total PUFA	53.02±2.88 ^a	46.62±1.98 ^b	15.60±0.25 ^d	11.43±0.38 ^e	34.75±1.47 ^c	40.85±1.89 ^b

Total PUFA includes n-6; n-3; 16;2; 16;3 and 16;4. Values below 0.5% for all tested diets are not included in this table. Mean±SD; n=6. nd: not detected.

1 **Figures**

2 **Figure 1.**

3 Average individual wet weight (mg) at the end of the four weeks feeding trial (mean±sd; n=6).

4 Superscripts indicate statistically significant differences.

5

6 **Figure 2.**

7 nMDS plot of the fatty acid profile of the six tested diets. Sample statistic (Global R)= 0.956;

8 Significance level of sample statistic= 0.01%; Number of permutations= 9999 (Random sample from

9 a large number); Number of permuted statistics greater than or equal to Global R= 0

10

11 **Figure 3.**

12 Abundance of the five main fatty acid groups from the experimental diets (mean±sd; n=6).

13 Superscripts indicate statistically significant differences.

14

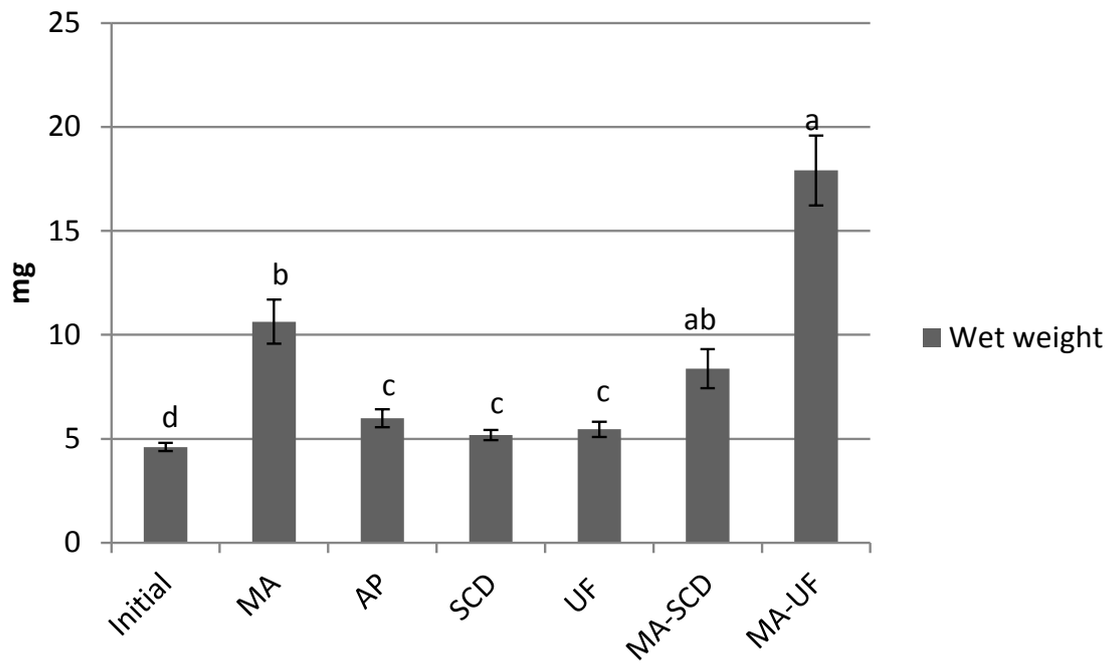
15 **Figure 4.**

16 Abundance of the main essential fatty acids (EPA, DHA and ARA) from the experimental diets

17 (mean±sd; n=6). Superscripts indicate statistically significant differences.

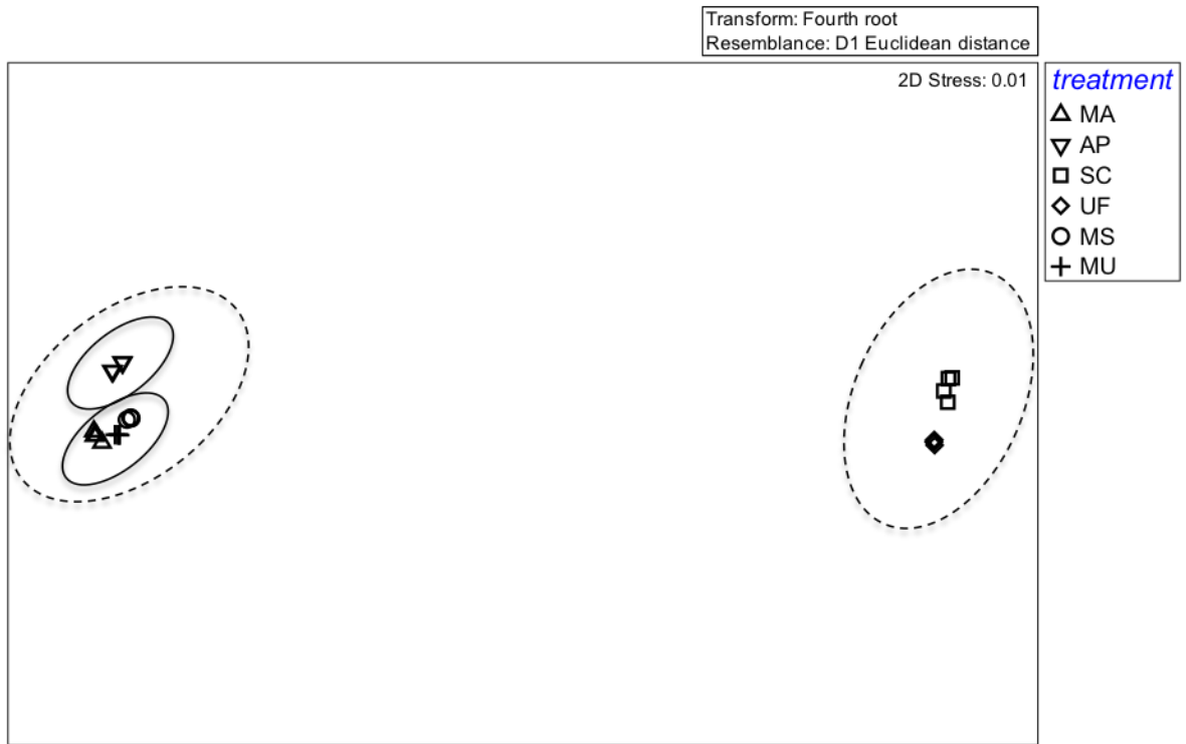
18

19 Figure 1



20

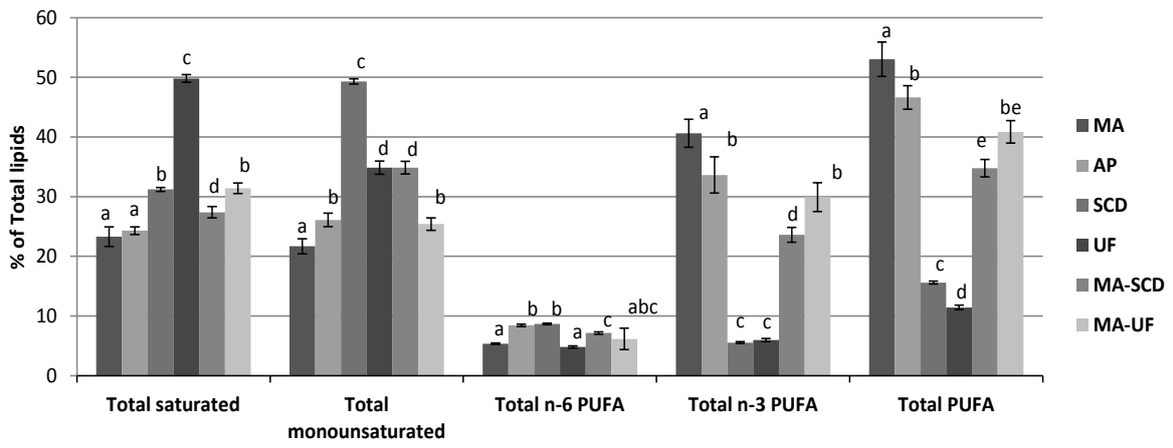
21 Figure 2



22

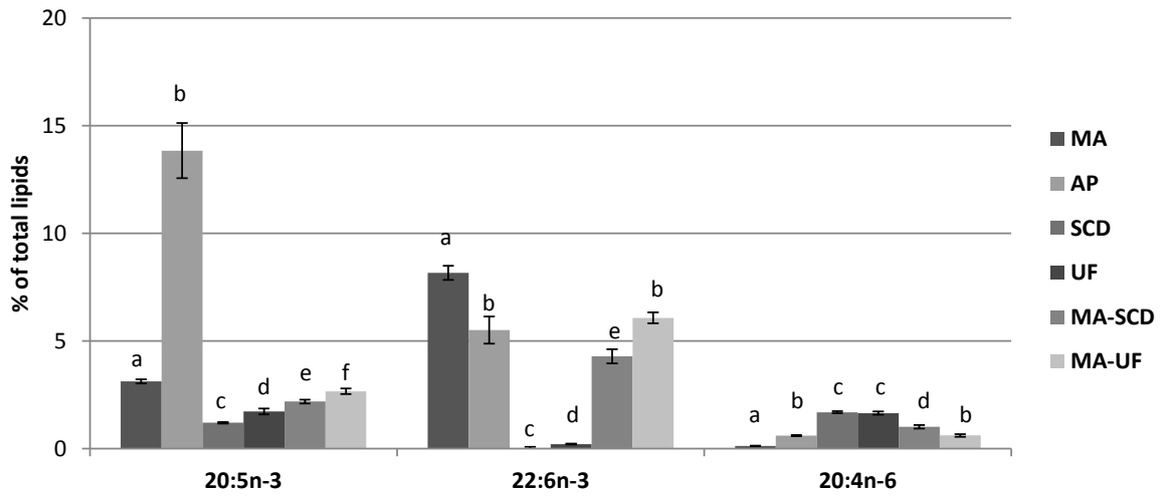
23

24 Figure 3



25

26 Figure 4



27

28