# Disentangling the counteracting effects of water content and carbon mass on zooplankton growth Kristian McConville<sup>1,2</sup>, Angus Atkinson<sup>1</sup>, Elaine S. Fileman<sup>1</sup>, John I. Spicer<sup>2</sup>, Andrew G. Hirst<sup>3,4</sup> 1 - Plymouth Marine Laboratory, Prospect Place, Plymouth, PL1 3DH, United Kingdom, krm@pml.ac.uk 2 - Marine Biology & Ecology Research Centre, School of Marine Science & Engineering, Plymouth University, Drake's Circus, Plymouth, PL4 8AA, United Kingdom 3 - School of Biological and Chemical Sciences, Queen Mary University of London, Mile End Road, London, E1 4NS, United Kingdom 4 - Centre for Ocean Life, National Institute for Aquatic Resources, Technical University of Denmark, Kavalergarden 6, Charlottenlund 2920, Denmark Keywords; water content, zooplankton, gelatinous, carbon percentage, growth

#### 27 ABSTRACT

Zooplankton vary widely in carbon percentage (carbon mass as a percentage of wet mass), but are 28 often described as either gelatinous or non-gelatinous. Here we update datasets of carbon 29 30 percentage and growth rate to investigate whether carbon percentage is a continuous trait, and whether its inclusion improves zooplankton growth models. We found that carbon percentage is 31 32 continuous, but that species are not distributed homogenously along this axis. To assess variability of this trait in situ, we investigated the distribution of biomass across the range of carbon percentage 33 for a zooplankton time series at station Plymouth L4. This showed separate biomass peaks for 34 gelatinous and crustacean taxa, however carbon percentage varied 8 fold within the gelatinous 35 36 group. Species with high carbon mass had lower carbon percentage, allowing separation of the counteracting effects of these two variables on growth rate. Specific growth rates,  $g(d^{-1})$  were 37 negatively related to carbon percentage and carbon mass, even in the gelatinous taxa alone, 38 39 suggesting that the trend is not driven by a categorical difference between these groups. The 40 addition of carbon percentage doubled the explanatory power of growth models based on mass 41 alone, demonstrating the benefits of considering carbon percentage as a continuous trait.

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# 43 INTRODUCTION

Gelatinous zooplankton are a phylogenetically broad and ecologically important group of taxa found throughout the world's oceans. Their prey range from bacteria to fish (Sutherland et al., 2010) and they exhibit an equally diverse range of life history strategies and body compositions. The high water content characteristic of this group can be expressed as carbon percentage (carbon mass as % of wet mass), with some taxa having carbon mass as low as 0.01% of their wet mass (Clarke et al., 1992; Harbison, 1992; Lucas et al., 2011; Kiørboe, 2013).

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Interest in gelatinous zooplankton is linked to a growing appreciation of their impact on pelagic ecosystems and human activities (Richardson et al., 2009; Purcell, 2012, Gibbons and Richardson, 2013). For example, the introduction of the ctenophore, *Mnemiopsis leidyi* to the Black Sea has had considerable financial implications for fisheries in the area (Shiganova and Bulgakova, 2000). Research on gelatinous zooplankton has grown apace with basic ecological interest in the

56 physiology, trophic ecology and bloom dynamics of this group (Møller and Riisgård, 2007;

57 Gemmell et al., 2013; Condon et al., 2013).

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Based on a compilation of zooplankton body composition, Kiørboe (Kiørboe, 2013) found that most 59 60 zooplankton species are either gelatinous ( $\sim 0.5\%$ ) or non-gelatinous (5-10%), with comparatively few intermediates. Indeed, much research has been directed toward comparing and contrasting 61 gelatinous versus non-gelatinous zooplankton. For example, compared to other planktonic animals, 62 gelatinous zooplankton have higher carbon mass-specific feeding rates (Hamner et al., 1975; 63 Acuña, 2001; Acuña et al., 2011), lower locomotion costs and higher specific growth rates (Hirst et 64 65 al., 2003; Pitt et al., 2013). Indeed, gelatinous taxa such as salps are amongst the fastest growing 66 metazoans (Bone, 1998).

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The use of a categorical approach to zooplankton body composition (i.e. gelatinous versus non-68 gelatinous) contrasts with the treatment of carbon mass (Peters, 1983), which is used as a 69 70 continuous variable in many models of growth (Hansen et al., 1997; Gillooly et al., 2002, Hirst et al. 2003). However, the carbon percentage of zooplankton species also varies widely, even among 71 gelatinous taxa (Molina-Ramirez et al. 2015). A recent review suggested that water content was 72 73 second only to body size in determining key aspects of the biology of zooplankton (Andersen et al., 2015b). So far, empirical models of zooplankton growth use equations that are specific to various 74 75 taxonomic groups (e.g. Hirst et al. 2003, Kiørboe & Hirst, 2014) and these equations have not yet been unified. As carbon mass and carbon percentage are both variable traits, it is important to 76 77 consider them together in empirical models of zooplankton growth. Furthermore, quantifying the 78 relationship between growth rate and carbon percentage may help to explain how carbon percentage 79 functions as an evolutionary trait, and, for example, why there are gelatinous representatives from 80 six phyla found in the plankton.

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In this study we have used both a meta-analyses approach and an in-situ time series of zooplankton from weekly sampling at the Plymouth L4 time series (Smyth et al. 2015). We had three objectives. The first was to quantify the degree of variability in carbon percentage both in "trait space" from the meta-analysis dataset and in a natural plankton assemblage, to gauge whether it was appropriate to treat water content as a continuous variable. The second aim was to investigate the degree of collinearity between carbon mass and carbon percentage, again both in a meta-assemblage and in

the L4 assemblage. Dependent on the outcome of these two objectives, the third aim was to
construct a simple empirical model of zooplankton growth that combines carbon mass and carbon
percentage.

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#### 92 *METHODS*

## 93 *Carbon percentage data*

94 Ratios of wet mass to carbon mass were combined from a series of recent compilations (Kiørboe, 95 2013;Pitt et al., 2013;Molina-Ramirez et al., 2015). The amalgamated dataset with their sources is 96 presented in Supplementary Information 1. Only concurrent measurements of carbon and wet mass 97 of the same individual were used to calculate carbon percentage.

The degree of tissue dilution of zooplankton taxa has been expressed previously as body carbon content (Molina-Ramirez et al., 2015). However to avoid confusion with carbon mass, throughout this paper it is referred to as "carbon percentage" (carbon mass as a percentage of wet mass). For our comparisons the levels of taxonomic organisation were selected based on functional diversity and body form (e.g. phylum for Chaetognatha, but orders Cydippida and Lobata).

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## 104 In situ analysis

To investigate how species biomass was distributed along the spectrum of carbon percentage an in 105 106 situ community, the L4 zooplankton time series (Western Channel Observatory, Plymouth) was used. The L4 sampling site is approximately 15km south-west of Plymouth and undergoes seasonal 107 stratification (Harris, 2010). Sampling at the L4 site consists of a pair of vertical hauls with a 200 108 um WP2 zooplankton net from 50 m to the surface (maximum depth 54m). The nets are retrieved at 109 20 cm s<sup>-1</sup> and are immediately fixed in 4% formaldehyde solution (Maud et al., 2015). The 110 zooplankton are then subsampled, counted and identified (Eloire et al. 2010). This zooplankton 111 abundance times series has high resolution both temporally (weekly sampling) and taxonomically, 112 with many taxa consistently identified to species level since 2009. To determine zooplankton 113 biomass, a total of 3780 individuals from the formalin-preserved catches at L4 taken throughout 114 2014 and 2015 were measured. From standard length measurements (e.g. cnidarian bell height or 115 diameter, copepod prosome length), length-carbon mass relationships from the literature were used 116 to estimate carbon masses per individual. These length measurements were then aggregated into 117 seasons, namely spring (March-May), summer (June-August), autumn (September-November) and 118

winter (December to February) to account for the high intraspecific variability in length observed at
L4 (Atkinson et al., 2015). This allowed us to derive season-specific mean carbon masses per
individual, which were multiplied by numerical densities to estimate biomass density (mg C m<sup>-3</sup>).
Previously measured, L4-specific seasonal values of individual carbon biomass were used, when
available (e.g. *Calanus helgolandicus;* Pond et al. 1996).

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Of the approximately 189 taxa recorded at L4, only 22 contributed more than 0.5% to the total biomass for all species. To examine how biomass was distributed across the spectrum of carbon percentage, these taxa were assigned to  $\log_2$  classes (0.1 - 0.2%, 0.2 - 0.4%, 0.4 - 0.8%, 0.8 - 1.6%, 1.6 - 3.2%, 3.2 - 6.4%, 6.4 - 12.8%, > 12.8%) using the carbon percentage data in Supplementary Information 1. The distribution of carbon biomass in each carbon percentage category across the seasons was then calculated.

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## 132 *Growth rate data*

Using the references from the appendices of Kiørboe and Hirst (Kiørboe and Hirst, 2014) as a starting point, zooplankton growth rate data were extracted from the original sources and augmented by searching the literature. All growth rate data used here are in Supplementary Information 2.

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To improve comparability of source data we restricted the meta-analysis to data from laboratory incubations with food available in high (assumed non-limiting) concentrations. By using only data collected under these conditions we suggest that the measurements are more directly comparable, with the observed patterns more likely to reflect the intrinsic biology of the species than external factors.

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Published growth rates are normally expressed either as increase in length or body mass over time. When organism size was expressed as length, published length-mass regressions were used to convert to body carbon mass (Hirst, 2012; Kiørboe and Hirst, 2014). To express growth rates in the terms commonly used for zooplankton (as an exponential rate; see Hirst and Forster 2013), the mass-specific growth rate, g (d<sup>-1</sup>) was determined as:

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$$g = (\ln M_t - \ln M_0)/d$$

150 where  $M_t$  is mass at time t,  $M_0$  is mass at the previous time point, and *d* is the time period between 151 the two measurements of mass (in days).

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Growth data were temperature-corrected to  $15^{\circ}$ C using a Q<sub>10</sub> of 2.8 (following Hansen et al., 1997; Kiørboe and Hirst, 2014). General linear models (GLMs) were constructed in R (R Core Team, 2014) to determine the relationships between growth rate, carbon percentage and carbon mass. To determine whether there was collinearity between the predictor variables we examined the condition indices for the variables in the model using the *colldiag* function in the *perturb* package in R (Hendrickx, 2012). A condition index of greater than 30 is considered large (Belsley et al., 1980) and suggests that the variable should be removed from the model.

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When growth data were available for a species but carbon percentage values were not, the latter was estimated using the mean value for the highest level of taxonomic relatedness available. For instance, if composition values for a species were not available, then the composition values for all other species within the genus were averaged and used as an estimate. The estimates were typically at the genus level but no lower relatedness than family (38% estimated at family level, primarily for copepods).

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## 168 *Growth rate analysis*

Four analyses were performed; the first two were based on mean and maximum growth rates for all 169 170 zooplankton taxa in the dataset, the second two as above but for the classical gelatinous taxa only (Cnidaria, Ctenophora and Thaliacea). Maximum growth values were defined as the highest 171 temperature-adjusted growth rate value available for each species. Issues of non-independence 172 173 between data were avoided by using single growth rate values per species per study. For illustrative purposes only (i.e. the plots in Fig. 4), we adjusted all growth rates to a fixed body carbon mass of 174 1 mg C after correcting to 15°C. This mass correction was performed assuming log<sub>10</sub> mass-specific 175 growth (g) scales against  $\log_{10}$  mass with a slope of -0.25 (Brown et al. 2004). 176

# 179 Variability in carbon percentage across the zooplankton

The range in body volume for two animals of equal carbon mass but at either end of the carbon 180 percentage spectrum is demonstrated in Figure 1. For the compiled dataset, the range in carbon 181 percentage extended over four orders of magnitude in zooplankton, from 0.01% in the lobate 182 183 ctenophore, Bathycyroe fosteri, to 19.02% in the copepod, Calanus hyperboreus (Fig. 1, 2a, Supplementary Information 1). The intervals between adjacent ranked species were small relative 184 185 to the range covered (Fig. 2a), suggesting that water content could be considered as a continuous variable. The largest interval between species coincided with the shift from the classic gelatinous 186 187 taxa to other zooplankton (i.e. from Thaliacea to Chaetognatha). However, this difference between species constituted a relatively small fraction of the total range (6.8%). In addition, there was 188 overlap of classic gelatinous and non-gelatinous groups. For example, some chaetognaths were 189 within the traditional gelatinous range (1.27% and 1.35% for *Pseudosagitta lyra* (as *P. scrippsae*) 190 and *Pseudosagitta* (as *Sagitta*) gazellae respectively), whereas one tunicate had a carbon percentage 191 which lay within the non-gelatinous range (3.87% for Doliolum denticulatum). This overlap of 192 taxonomic groups was extensive across the spectrum of water content, as can be seen by the mixing 193 194 of colour across Fig. 2. This was particularly the case among the Ctenophora and Thaliacea with the 195 range of both taxa approaching two orders of magnitude in carbon percentage.

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The wide variation in body carbon percentage observed at a species level in Fig. 1a is also 197 summarised at the broader taxon level in Fig. 2b. Median values for groups do loosely cluster into 198 gelatinous and non-gelatinous taxa following the bimodal distribution of species suggested by 199 200 Kiørboe (Kiørboe, 2013). The ranges of all adjacent taxa (excluding lobate ctenophores) overlapped, with Thaliacea and Chaetognatha bridging the gap between the classical gelatinous and 201 non-gelatinous taxa. The variability within groups was greater for gelatinous taxa, with the greatest 202 203 range in the scyphomedusae, closely followed by the thaliaceans. The gelatinous taxa sort into their 204 respective phyla when ranked (i.e. Lobata, Nuda, Cydippida for the Ctenophora, then 205 Hydromedusae and Scyphomedusae for Cnidaria) suggesting that taxa within phyla are on average 206 more similar to each other than with other phyla.

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In the natural assemblage sampled at the Plymouth L4 site (Figure 3) we have an alternative picture, namely how biomass is distributed along this spectrum of carbon percentage. At L4, biomass is distributed bimodally. The biomass is primarily concentrated in the categories that are either highly

gelatinous (carbon mass 0.1 - 0.8% of wet mass) or non-gelatinous (6.4 -> 12.8\%) However, there 211 is considerable variability within the carbon percentage categories, as some gelatinous taxa are as 212 213 much as 8 times larger in wet mass for the same carbon mass as others. The biomass in the intermediate categories (0.8 - 1.6%) and 1.6 - 3.2%) was very low and below our threshold for 214 215 inclusion. This area of the spectrum is populated by thaliaceans and large rhizostome scyphomedusae, which are either not commonly recorded at L4 (thaliaceans) or are rarely or poorly 216 sampled by the 0.57 cm diameter nets used. Gelatinous taxa comprise a greater proportion of 217 218 biomass in summer than the other seasons. In winter, chaetograths (3.56%) have similar total biomass to the dominant copepods. There is also a broad trend of increasing carbon percentage 219 through the year within the gelatinous taxa. In spring, the cydippids (the most gelatinous group 220 frequently encountered at L4) are dominant, followed by Nuda (Beroe) in summer and finally 221 hydromedusae and siphonophores in autumn. 222

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# 224 *Relationship between carbon mass and carbon percentage*

225 There were negative relationships between carbon mass and carbon percentage, both in the metadataset (Fig. 4a) and in the in situ dataset (Fig.4b). While the more gelatinous taxa tended to have 226 higher carbon mass there was considerable variability, with some organisms of similar carbon mass 227 differing 100-fold in carbon percentage (Fig. 4). To ensure that collinearity was not influencing the 228 growth model the condition indices for the variables were inspected. The highest condition index 229 observed was 3.05, lower than the threshold of 30 suggested by Belsley (Belsley et al., 1980) 230 confirming that carbon mass and carbon percentage can be used in combination in models of 231 zooplankton growth. As gelatinous and small organisms tend to grow fastest, the tendency for more 232 gelatinous taxa to have higher carbon mass underlines the need to include both as covariates in our 233 growth model. 234

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## 236 *Relationship between carbon percentage and growth rate*

We first conducted GLMs on the subset of data comprising the classical gelatinous taxa alone. These showed that mean growth rate declined with increasing mass and increasing body carbon percentage. The GLMs on the whole dataset established that  $log_{10}$  mass-specific mean and maximum growth rate was significantly correlated with both  $log_{10}$  carbon mass and  $log_{10}$  body carbon percentage (Fig. 5, Table I). As expected, there was a negative relationship between  $log_{10}$ mass-specific growth rate (g), and  $log_{10}$  carbon mass, in line with the results of Kiørboe and Hirst (Kiørboe and Hirst, 2014). In the analyses of all zooplankton taxa, mean and maximum growth ratedecreased with increasing carbon mass and carbon percentage.

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In all analyses, the addition of body carbon percentage to models of growth based on carbon mass 246 247 alone increased the explanatory power (Table II). The second order Akaike criterion, AICc, (Burnham and Anderson, 2002) was lower in the model including water content in all analyses, 248 supporting the inclusion of this factor in analyses of zooplankton growth. In the maximum analysis 249 including all taxa, Akaike weights  $(\omega_i)$  were approximately 10 times higher in the models including 250 251 body carbon percentage (mass  $\omega_i = 0.08$ , mass + carbon percentage  $\omega_i = 0.92$ ). This suggests that these models were significantly better than models based on mass alone (Royall, 1997). A similar 252 253 pattern was observed in the analysis of maximum growth rates of the gelatinous taxa however it was not observed for mean growth rates (mass  $\omega_i = 0.02$ , mass + GI  $\omega_i = 0.98$ ). 254

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### 256 DISCUSSION

Our study provides strong support for: body carbon percentage being a continuous trait, for a negative relationship between body carbon percentage and growth rate, and for considerable increases in model predictive power as a result of inclusion of this trait for zooplankton. Below we discuss the implications of each of these findings in turn.

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Kiørboe (Kiørboe, 2013) demonstrated that if zooplankton are arranged in a frequency distribution 262 based on body composition, that most taxa are either gelatinous (carbon mass is ~0.5% of wet mass) 263 264 or non-gelatinous (~5-10%), with little overlap. Our study would appear to contradict this, since we found a fairly continuous distribution of carbon percentage. However, this does not conflict with the 265 findings of Kiørboe (Kiørboe, 2013), since in that study it was emphasised that most taxa are either 266 highly gelatinous or non-gelatinous. Rather, we highlight that, while the most species fall into one 267 of these two groups, there is considerable variability in carbon percentage within each group and 268 there are representatives across much of this spectrum. The distribution of zooplankton biomass at 269 L4 supports both of these views. Biomass is clustered at either end of the spectrum as described 270 previously, and this could suggest that the fitness landscape for this trait favours extremes. 271 However, at either end of the spectrum there is considerable variability. The traditional gelatinous 272 273 group alone spans an 8-fold range in carbon percentage, with implications for growth rate. For

example, there is a trend of increasing carbon percentage among the gelatinous zooplankton through
the year, with cydippids being replaced by beroids in summer and finally by hydromedusae and
siphonophores in autumn.

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In the meta-analysis compilation, the largest interval occurs between taxa typically considered as 278 gelatinous and intermediate, between the pelagic tunicate, Thalia (as Salpa) democratica (1.6 % 279 body carbon percentage) and a chaetognath, Eukrohnia hamata (2.7 % body carbon percentage. 280 Molina-Ramirez et al. (Molina-Ramirez et al. 2015) stressed that considerable variation in carbon 281 percentage existed even within the classic gelatinous taxa (Cnidaria, Ctenophora and Tunicata). Our 282 results are in agreement, albeit with even higher degree of variability (at 350-fold). Taken together, 283 284 the relatively small interval between values for gelatinous and non-gelatinous species and the high variability observed within the gelatinous taxa suggest that growth models can indeed incorporate 285 286 carbon percentage as a continuous trait.

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When  $\log_{10}$  mass-specific growth rate was regressed against  $\log_{10}$  body carbon percentage as a continuous variable, a negative relationship was observed. Crucially, the pattern persisted when considering the gelatinous taxa alone (Table II). The existence of the relationship among the gelatinous taxa alone, is important as this demonstrates that the relationship is not due to a categorical difference between gelatinous organisms and non-gelatinous organisms.

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One potential mechanism that could explain the relationship between body carbon percentage and 294 growth rate is enhanced feeding rate (Acuña et al., 2011). These authors suggested that the large 295 296 dilute bodies of gelatinous zooplankton facilitate higher carbon-specific feeding rates than other zooplankton taxa of the same carbon mass. If this increased feeding rate drives faster growth, then 297 this might explain the relationship of increasing growth rate with decreasing carbon percentage (see 298 Fig. 2). As many gelatinous taxa are filter or ambush feeders that rely on capture surfaces to feed, 299 300 assuming that feeding rate scales with surface area, then we may expect the scaling exponent between surface area and body carbon percentage to match the exponent for growth rate and body 301 302 carbon percentage. To investigate this we used a simple geometric calculation. Assuming 303 isomorphic growth, surface area (SA) scales with body volume with a power of 0.67. By altering 304 degree of gelatinousness for a fixed amount of body carbon, SA then scales with carbon percentage with a power of -0.67. Hence, with an assumption that growth rate is a fixed proportion of feeding 305

rate, this would give the same slope of -0.67 for  $\log_{10}$  mass-specific growth versus  $\log_{10}$  carbon percentage (Fig. 2). The exponents that we determined empirically across the various zooplankton taxa are less steeply negative than -0.67 (at -0.18 and -0.16 for mean and maximum respectively), i.e. increasingly gelatinous organisms increase their growth rate less rapidly than these surface considerations would predict. This could indicate a potential feeding inefficiency associated with decreasing carbon percentage or that factors additional to surface area may also be important.

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In common with Ikeda (Ikeda, 2014), we found that species with larger total carbon masses also 313 tended to be more watery. Furthermore, as the larger organisms are typically more watery the 314 effects of carbon mass and carbon percentage tend to counteract, underscoring the need to include 315 these variables together in order to better predict growth. Molina-Ramirez et al. (Molina-Ramirez et 316 al., 2015) found a similar result for tunicates but found that body carbon percentage was invariant 317 318 with increasing mass for cnidarians and ctenophores. The authors suggested that this might be due to differences between internal filter feeding in tunicates and external ambush or cruise feeding in 319 320 the other groups. It has been suggested that feeding modes decrease in efficiency with increasing size (Kiørboe et al. 2011), so high water content may help to mitigate this decrease in efficiency and 321 322 maintain relatively higher carbon specific feeding rate at large carbon masses. This is supported by the findings of Acuña et al. (Acuña et al., 2011), suggesting that gelatinous plankton had higher 323 324 carbon-specific feeding rates than other zooplankton of a similar carbon mass. Together with higher growth rates, these factors could help to explain how gelatinous zooplankton are capable of forming 325 such high localised increases in species biomass (blooms). 326

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While the increase in capture surface area and associated feeding and growth rates is one potential 328 329 advantage of the gelatinous body form, there are other implications. There are potential negative implications also, especially with regard to limited swimming speed and escape responses. While 330 medusae have potential defences in the form of nematocysts, many gelatinous taxa such as 331 ctenophores do not, and may have limited ability to escape from potential predators as a result of 332 333 their large dilute bodies (Acuña, et al. 2011). Understanding why some taxa are gelatinous is not 334 always straightforward. The most gelatinous mollusc in this analysis is Clione limacina, a gymnosome predator that feeds on almost exclusively on *Limacina helicina*. Clione does not rely on 335 large capture surfaces or on generating a feeding current as it ambushes individual, relatively large 336 prey items. In this case, water content does not appear to be a derived trait to increase body volume 337 relative to carbon for feeding, suggesting that this may not be the only driver of high water content 338

in zooplankton. It has been suggested that potential other causes include physical or ecological
factors such as transparency to impair visual predation (Hamner et al., 1975) or the efficiency of
neutral buoyancy (Kiørboe, 2013). Together these factors may help to explain why semi-gelatinous
bodies are observed in at least six major planktonic phyla (Cnidaria, Ctenophora, Chordata,
Annelida, Chaetognatha, Mollusca, see Supplementary Information 1).

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# 345 CONCLUSIONS

Body size is often described as a master-trait, and is frequently used as the sole intrinsic variable in 346 empirical and simulation models involving zooplankton growth (Kiørboe and Hirst, 2014, Anderson 347 et al., 2015a). But what do we mean by "body size"? Carbon mass is often used as the unit for size, 348 but both our meta-analysis and the real assemblage data show that carbon percentage also varies 349 greatly. It may even vary negatively with carbon mass, levering an opposing effect on growth. We 350 351 argue that carbon mass and carbon percentage are both key traits, both are intrinsic to the zooplankton and since they are possible to estimate, then we should disentangle their separate 352 353 effects in a unified growth model. By including carbon percentage to models of growth based on carbon mass alone, we substantially increased their explanatory power, with smaller body masses 354 355 and lower body carbon percentages leading to higher specific growth rates. Building on the work of previous publications (Kiørboe, 2013, Pitt et al., 2013, Molina-Ramirez et al., 2015) we provide a 356 carbon percentage dataset in Supplementary Table 1. By using these source data alongside carbon 357 masses, the maximum growth rate equation in Table 1 may then be used as a starting point to 358 estimate growth rates attainable by zooplankton. 359

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Alongside the "size" based simplifications used for modelling, there has also been an increase in 361 362 "trait-based" modelling in which categorical variables or functional groups are allowed to vary 363 continuously. A purpose of this paper is to allow water content also to be used as a continuous trait; to facilitate its inclusion alongside carbon mass and other traits such as feeding mode (Litchman 364 365 2013; Andersen et al. 2015a; Hérbert et al., 2016). Since we found that growth rate depended on carbon percentage even among the gelatinous taxa alone, we hope that considering and modelling 366 water content as a continuous trait will reveal the ecological and evolutionary factors that influence 367 the water content of zooplankton. 368

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490 TABLE AND FIGURE LEGENDS:
491 Figure 1. Comparison of the relative carbon (black) and wet masses (grey) of *Calanus hyperboreus*492 (left, carbon percentage = 19.02%) and *Bathycyroe fosteri* (right, carbon percentage = 0.01%). The

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Figure 2. (a) Zooplankton species ranked according to their carbon percentage (CM%WM;log<sub>10</sub> scale), each horizontal bar represents a single species. Colours indicate taxonomic groups as detailed in the legend. (b) Zooplankton taxonomic groups ranked according to their carbon mass (as % of wet mass; log<sub>10</sub>scale). Boxes indicate median, lower and upper quartiles with whiskers showing the range. (Vertical lines at 0.5 and 5 CM%WM represent the composition of the gelatinous and non-gelatinous taxa defined by Kiørboe 2013).

relative area of each shade is scaled as volume so the silhouettes are representative of true size.

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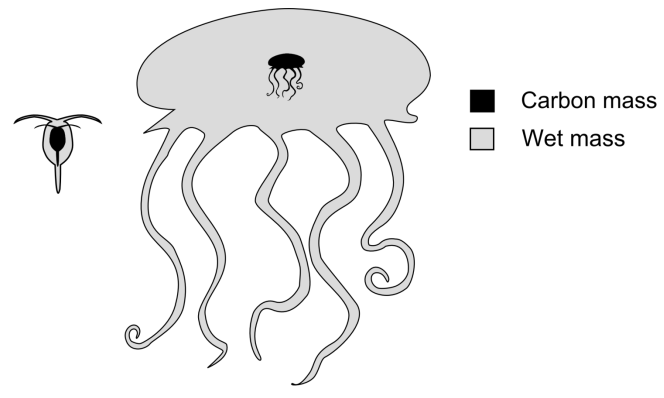
Figure 3. Distribution of carbon biomass (mg C m<sup>-3</sup>) between  $\log_2$  carbon percentage (CM%WM) categories through spring, summer, autumn and winter (2009-2015) at the L4 sampling site, Western Channel Observatory, Plymouth. The same colour coding of taxa is used as in Fig. 1 – see legend. \* - Biomass value for the category 0.4 – 0.8 exceeds the scale in summer (34.4 mg C m<sup>-3</sup>) as a result of 7 high abundance observations of *Beroe spp*. (of total 318 samples). Upper limit of biomass scale in winter is 5 mg C m<sup>3</sup>.

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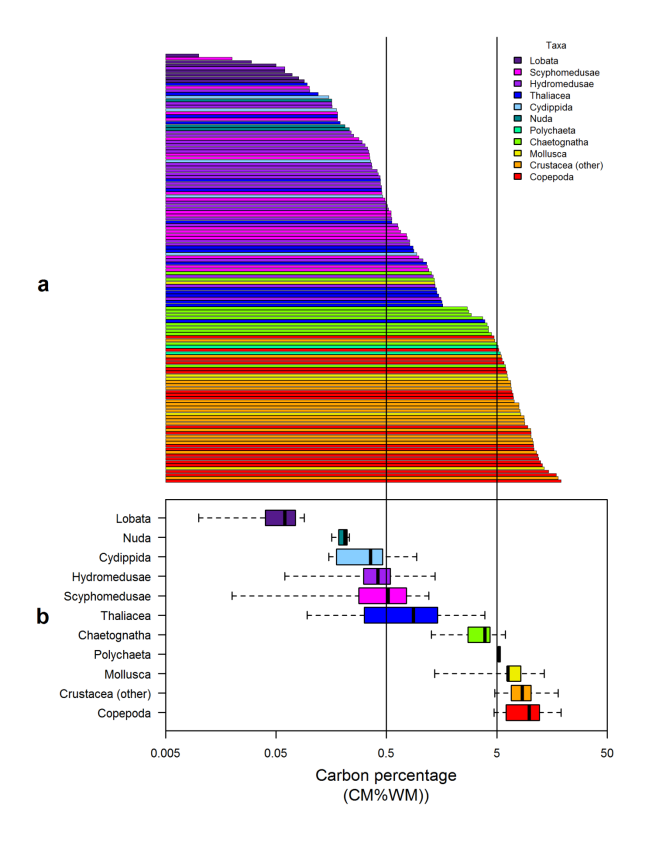
Figure 4. Carbon percentage (CM%WM) as a function of carbon mass (mg) for the meta-analysis dataset (A, log carbon percentage =  $-0.26 \times \log$  carbon mass -0.18, p = 0.0001, R<sup>2</sup> = 0.21, df = 60) and the L4 assemblage (B, log carbon percentage =  $-0.34 \times \log$  carbon mass -1.1, p = 0.0026, R<sup>2</sup> = , df = 20). Taxonomic groups coloured as indicated in the legends.

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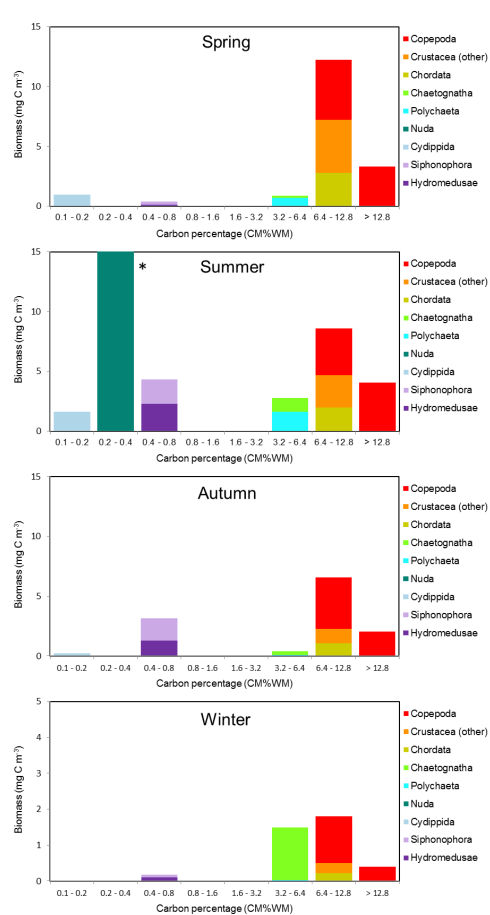
514	Figure 5. Specific growth rate, $g(d^{-1})$ as a function of body carbon percentage (CM%WM). Growth
515	values were temperature-adjusted to 15°C, mass adjusted to 1 mg C and then averaged for each
516	species in each study. (a) mean mass-specific growth rate values for each species in each study and
517	(b) maximum specific growth rate values for each species.



540 Fig 1

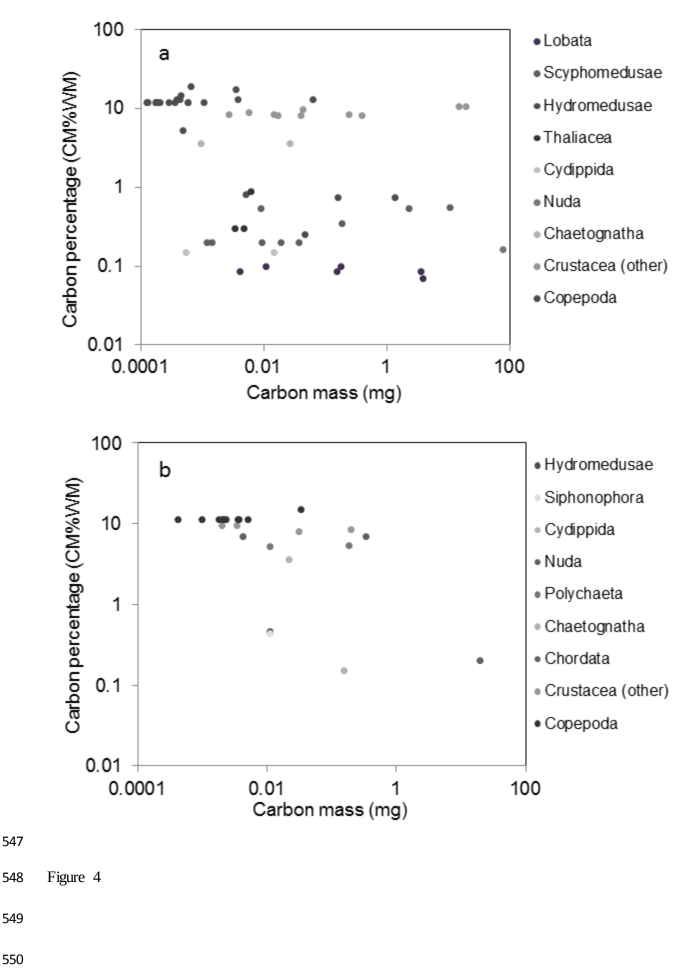


542 Fig 2





546 Fig 3





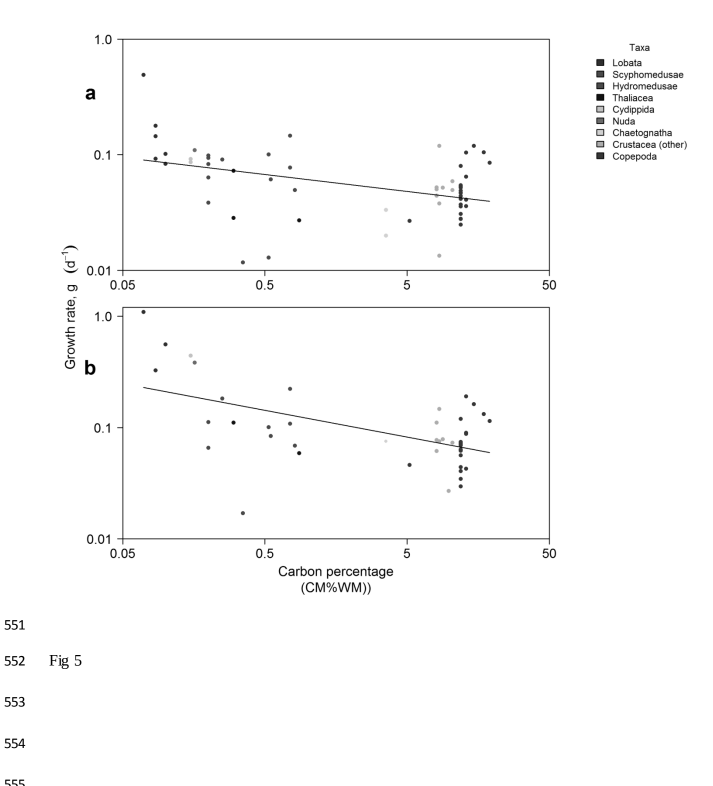


Table I. General linear models predicting  $\log_{10}$  mean specific and  $\log_{10}$  maximum specific growth rate, g (d<sup>-1</sup>), as a function of both  $\log_{10}$  carbon mass (mg) and  $\log_{10}$  body carbon percentage (100\*(CM/WM)). All models pertain to growth rate data that were first Q<sub>10</sub>-adjusted to 15°C.

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Table II. Changes to measures of explanatory power of models of growth based solely on carbon mass when body carbon percentage (CC) was added as a factor. AICc is the corrected Akaike information criterion,  $\Delta_i$  is the AIC difference, and  $\omega_i$  is the Akaike weight. Models with Akaike weight values 10 times greater than that of the other models being compared are considered statistically significant as optimal models (mass + GI for mean and max all zooplankton and max gelatinous taxa only). All models pertain to growth data that were first Q<sub>10</sub>-adjusted to T = 15°C.

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Group		Factor	df	р	Slope	Intercept	Adj R <sup>2</sup>
All zooplankton	Mean growth rate,g	$log_{10}$ carbon mass $log_{10}$ carbon percentage	58	<0.0001 0.036	-0.17 -0.18	-1.12	0.43
	max growth rate,g	$log_{10}$ carbon mass $log_{10}$ carbon percentage	42	<0.0001 0.013	-0.16 -0.16	-0.81	0.31
Gelatinous taxa only	mean growth rate,g	$log_{10}$ carbon mass $log_{10}$ carbon percentage	22	0.027 0.038	-0.19 -0.17	-1.18	0.33
	max growth rate,g	$log_{10}$ carbon mass $log_{10}$ carbon percentage	13	0.011 0.018	-0.16 -0.72	-1.15	0.42

Table I

C	n	n
ю	u	U

000	Group	g	$\mathbf{R}^2$		AICc		$\Delta_{ m i}$	ω <sub>i</sub>	
601			Mass	Mass + CC	Mass	Mass + CC		Mass	Mass + CC
	All	Mean	0.39	0.43	18.63	16.67	2.47	0.19	0.81
602	zooplankton	max	0.22	0.31	21.99	17.57	4.42	0.076	0.92
	Gelatinous	mean	0.33	0.33	18.51	19.96	1.44	0.54	0.46
603	taxa only	max	0.09	0.42	21.55	16.26	5.29	0.019	0.98

