

# Larval growth of the polychaete Arenicola marina under different temperature and food conditions: consequences on bioenergetic models

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1	Larval growth of the polychaete Arenicola marina under different
2	temperature and food conditions: consequences on bioenergetic models
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15	Lay summary
16	Biphasic growth models of Arenicola marina larvae highlight an optimal temperature at 15°C
17	with a lower temperature tolerance range compared to juveniles and adults. We claim that two
18	sets of Arrhenius temperatures depending on the life-history stages should be implemented
19	when using an abj-DEB model in this species.
20	

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#### 22 Abstract

23 Arenicola marina, a marine benthic polychaete, is widespread on sandy beaches in Europe and considered as an ecosystem engineer despite commonly used as bait by fishermen. Data 24 25 regarding the bioenergetics of the lugworm larval stages are still incomplete. Trochophore is initially lecithotroph and then becomes planktotroph while growing as metatrochophore on 26 27 subtidal area, a quite stable daily temperature environment compared to the foreshore, where 28 juveniles and adult live, with daily temperature fluctuating up to 15°C. These discrepancies in 29 temperature ranges may influence the temperature corrections that control metabolic rates during the life cycle of A. marina. We carried out laboratory experiments in microcosms by 30 31 inducing artificial spawning of lugworms, and then undertaken in vitro fertilization to obtain embryos and finally to follow, the larval development up to 10 segments with chaetae for 50 32 days under three temperature conditions (13°C, 15°C and 17°C) and two food conditions 33 34 ('fed' and 'non-fed'). The first feeding ('birth') of A. marina larvae was deciphered anatomically for a size between ~  $450-500 \mu$ m and described at 17 days post-fertilization for 35 larvae reared at 15°C and 17°C. Using a biphasic model with a von Bertalanffy growth before 36 'birth' and, an exponential growth after 'birth', among the three temperature treatments, the 37 15°C condition exhibited the best larval performance. Temperature corrections (TC) based on 38 39 embryonic and larval metabolic rates gave an Arrhenius temperature of ~ 6661 K and a higher boundary temperature tolerance range of ~ 294.5 K. Both temperature values differ from 40 those calculated from TC based mostly on juvenile and adult metabolic rates. We claim to use 41 42 two sets of Arrhenius temperatures according to the life-history stages of A. marina while using Dynamic Energy Budget model. This model was developed initially in order to manage 43 the conservation of the lugworm species. 44

Key words: Growth rate; lugworm; polychaetes; biphasic model; DEB model; larval stages;Arrhenius temperatures.

#### 47 Introduction

48 Polychaeta are mainly marine metazoans and represent significant part of the benthic biomass (Grémare et al., 1998). They play a major role in the functioning of benthic ecosystems and 49 serve as bio-indicators of the marine environment health status (Giangrande et al., 2005; 50 Sivadas et al., 2010). Polychaetes have a market value in fisheries where they are used as bait 51 by fishermen (Watson et al., 2017). In aquaculture, polychaetes may be used either as food 52 53 supplements due to their high nutritional value for cultured aquatic species (Pairohakul *et al.*, 2021) or by their abilities in waste depollution in integrated aquaculture (Jansen et al., 2019; 54 Jerónimo et al., 2020). Finally, studies have shown the therapeutic interest which certain 55 species of polychaetes may have for applications in human health (Kuijk & van Die, 2010; 56 Singh et al., 2014). For all these reasons, the breeding and marketing of polychaetes are of 57 growing interest and are current issues (Olive, 1993; Olive, 1994; Micael et al., 2016). This 58 59 attractiveness causes intensive harvesting of these species, mainly on foreshore, which results in an alteration of the environment and therefore has a deleterious effect on benthic 60 ecosystems (Beukema, 1989; Clarke et al., 2017). Moreover, this overexploitation of the 61 resource endangers the survival of some species of polychaetes (Cole et al., 2018; De Cubber 62 et al., 2018). To overcome these issues, some countries have implemented regulations that 63 64 aimed at regulating the rate of withdrawals over the years in sensitive areas e.g. in Portugal (Xenarios et al., 2018), in USA (Sypitowski et al., 2009), in Australia (Cole et al., 2018) and 65 in UK (Watson et al., 2015). Another way to avoiding the depopulation of polychaetes is to 66 develop the domestication of species of high economic interest; thus, farms of Atilla virens 67 (Olive, 1999; Sustainable Feeds Ltd<sup>TM</sup>), Arenicola marina (Olive et al., 2001; Hemarina 68 Ltd<sup>TM</sup>), Arenicola defodiens (Olive et al., 2001), Hediste diversicolor (Bischoff et al., 2009), 69 Diopatra aciculata (Safarik et al., 2006), Perinereis cf. nuntia (Poltana et al., 2007) and 70 Perinereis helleri (Palmer et al., 2016) have emerged. However, a complete knowledge of the 71

physiology of these polychaetes and in particular of the early stages of their development, is
necessary to carry out these conservation and cultivation projects.

The lugworm Arenicola marina (Linnaeus, 1758) is one of the most used bait for 74 professional and recreational fishing in Western Europe, where it is intensively harvested 75 from the Arctic to the Mediterranean (Watson et al., 2017; De Cubber et al., 2018). Moreover, 76 the strong affinity of its hemoglobin for oxygen has led to the production of this worm for 77 78 therapeutic uses in human health, whether as an organ preservative during transplants but also 79 as a possible blood substitute (Rousselot et al., 2006; Batool et al., 2021). A. marina lives in 5 to 40 cm deep U-shaped burrows in soft foreshore sediments in the intertidal area, from 80 81 mediolittoral to infralittoral (De Cubber et al., 2020). The life cycle of A. marina has been described in details (Newell, 1948; Newell, 1949; Farke & Berghuis, 1979a,b; Reise et al., 82 2001; De Cubber et al., 2019). Juveniles and adults live in burrows, where they swallow the 83 84 sediment at the surface being psammivorous. Lugworms may move backwards in the burrow, where they expulse their faeces by their pygidium that forms a characteristic sand-pile called 85 castings. Breeding season occurs in autumn where lugworm's population have annual 86 epidemic spawning of few days (Watson et al., 2000). Females spawn their oocytes within the 87 gallery, while males release sperm puddles on to the sediment surface that will be diluted by 88 89 the incoming tide and then drawn into female's gallery by pumping. Fertilization takes place inside the gallery (Williams et al., 1997) where embryos remain until hatching at the 90 trochophore larval stage. Trochophores and then metatrochophores are lecithotrophic larvae 91 92 dispersing several days (depending on temperature) in the water column until temporally (few 93 months) settling on subtidal marine habitats such as macroalgae or mussel beds (Farke & Berghuis, 1979a,b; De Cubber et al., 2019). During that first settling period, the first food 94 intake ('birth'; Dynamic Energy Budget (DEB) theory see after; Kooijman, 2010) occurs, 95 where larvae will live in a mucus tube but going out of their tube to collect organic matter or 96

phytoplankton. Larvae will develop segments with chaetae called setigers (up to 19 setigers) 97 98 until the completion of metamorphosis that could last up to 7 months (Farke & Berghuis, 1979a,b; De Cubber et al., 2019). When metamorphosis will be completed, a second phase of 99 100 dispersal will occur into the water column allowing post-larval stages to reach the foreshore. 101 These post-larvae will then settle on high part of the shore, burrowing themselves and 102 becoming a psammivorous juvenile as the adults. While growing and acquiring maturity to 103 become an adult, lugworms will migrate lower on the shore (De Cubber *et al.*, 2020). 104 Although the overall functioning of the life cycle is known, knowledge was still poor regarding the fine tune of the larval stage development of A. marina on the subtidal area 105 106 (Newell, 1948; Farke & Berghuis, 1979a,b; De Cubber et al., 2019).

To overcome this, and thus have a better knowledge of the different life stages of this 107 species, an abj-DEB model was developed by De Cubber et al. (2019). Indeed, DEB models 108 109 allow to predict the physiological processes (such as growth, assimilation, respiration, reproduction) of a species across its whole life cycle according to environmental conditions 110 111 (such as food availability and temperature) (Kooijman, 2010). When applying DEB theory (Kooijman, 2010), abj-DEB model (Margues et al., 2018) differs from a standard DEB model 112 by adding an extra juvenile life stage that takes place between the first feeding of the larval 113 114 stage (birth, 'b') to the end of the metamorphosis ('j') at the post-larval stage, where metabolic acceleration  $(s_M)$  occurs leading to an exponential growth of the individual 115 (Kooijman, 2014), compared to a classical von Bertalanffy growth before 'birth' and from the 116 juvenile to adult stages (Kooijman, 2010). However, data used for the abj-DEB model 117 118 developed by De Cubber et al. (2019) were not supported by data for the early-life stages 119 between the trochophore and the post-larval stages despite some predictions of age and length were obtained by simulation. No experimental studies have described so far, the early larval 120 121 stages of A. marina into details regarding the age versus length according to temperature and

food level. Most studies were focused on fertilization success and temperature effect during embryogenesis (prior the trochophore stage) at a stage that embryos still live into the female gallery on the foreshore (Lewis *et al.*, 2002; Watson *et al.*, 1998). In addition, abiotic factors such as temperature and food availability have not been tested in the laboratory to determine their effect on larval growth and development.

Thus, we carried out an experimental study in laboratory in order to deepen our 127 128 knowledge on the influence of temperature and food on the physiology of the larval stages of A. marina. The study aimed to determine precisely when the first feeding ('the birth': in DEB 129 theory) occurs (age at 'birth' and length at 'birth') in order to describe the biphasic growth 130 131 before and after 'birth' according to different temperature and food conditions. The second goal of this paper was to decipher if there was a difference into the thermal tolerance during 132 the life cycle of A. marina between different life stages as larval stages live in the subtidal 133 areas, a quite stable daily temperature, whereas juveniles and adults live in the intertidal areas, 134 where daily temperature can fluctuate up to 15 °C. These discrepancies in temperature ranges 135 136 in these two marine habitats may result in different sets of Arrhenius temperatures (Kooijman, 137 2010) that control metabolic rates of the lugworms according to its life stage. Overall data could be used to improve the existing abj-DEB model that has been developed initially in 138 139 order to help stakeholders to make decision for preserving A. marina in areas with high anthropogenic pressure or to improve the farming of this species in aquaculture. 140

141

#### 142 Materials and methods

143 Study area and sampling

144 For the need of our experiment, 180 adult lugworms were collected at Wimereux (50°46'N,

145 1°36'E), located on the Eastern English Channel, part of a marine protected area (MPA)

146 created in 2012. The coastline is principally composed of sandy beaches as well as rocky

shores mainly colonized by algae and mussels on the intertidal and subtidal areas (Rolet *et al.*,

148 2015). In this MPA, adult population of Arenicola marina are found on the high and mid-

shore (De Cubber *et al.*, 2018). From September  $2^{nd}$  to  $16^{th}$  2019, 180 adults of A. marina

were sampled in total, using a shovel and a bait pump (Decathlon ltd.) on the mid-shore at lowtide.

152

153 Broodstock selection and maintenance

At the Wimereux Marine Station, collected lugworms (n = 180) were maintained in a 300 L 154 tank with a continuous seawater flow (300 L.h<sup>-1</sup>), placed on a thermostatically controlled 155 room (15°C). A continuous flow of water mixed the tank for 24 hours in order to clean the 156 worms by removing sand and micro-algae residues. Then, to assess the reproductive status of 157 158 each worm, biopsies of the coelomic fluid were performed using a sterile hypodermic syringe on anaesthetized individuals Arenicola marina in three successive ethanol solution (1%, 2.5% 159 160 and 5%) in twice-filtered seawater solutions (TFSW, 0.45 µm and 0.2 µm) (Gaudron & 161 Bentley, 2002). Observations using an optic microscope (Motic® BA210) allowed to establish the state of maturity of the gametes and to differentiate the sexes. After sex 162 determination, males and females were separated and kept in two different tanks with 163 164 continuous seawater flow. While maintaining the lugworms, regular gametes observations using the optic microscope, were carried out randomly on biopsies of five males and five 165 166 females in order to estimate the reproductive status of each individual. For females, 30 random oocytes were measured using the optic microscope equipped with Motic Image Plus© 167 3.0 software. Female gametes were estimated to be ready for fertilization when mean oocytes 168 169 diameter was at 180 µm (Watson et al., 1998). For male gametes, maturity was fixed when 80 % rate of spermatocytes were in the morula stage (Dillon & Howie, 1997). 170

#### 172 Spawning induction, artificial fertilization

Five females and five males with the most mature gametes were selected as broodstock for 173 artificial fertilization. Lugworms were washed with autoclaved TFSW and then placed in 174 175 individual tanks (15.0 x 8.0 x 10.0 cm) filled with 1 L of TFSW. Each selected female was injected with two prostomial homogenates (Howie, 1961) and kept for 24 to 48 hours in an 176 individual tank at 15°C until the release of the oocytes. After spawning, females were 177 178 removed from their tanks and oocytes were collected with a 63  $\mu$ m-mesh. Then, female 179 gametes were washed twice with TFSW and stored in 5 mL microtubes at 4°C. Just after the release of oocytes, each male was injected with two prostomial homogenate 180 181 (Pacey & Bentley, 1992) and monitored until gametes release. After ejection by male's nephridiopores, 'dry' sperm was collected immediately with a micropipette and placed in 1 182 mL microtubes on ice (Williams et al., 1997). Male gametes were counted using a Neubauer 183 counting chamber (Sigma ltd.) under the optic microscope. Before the artificial fertilization, 184 females (n = 5) and males (n = 5) gametes were pooled together to increase fertilization 185 success. Approximately  $10^6$  oocytes were mixed with a concentration of  $10^4$  sperm per egg in 186 a 2 L autoclaved glass container filled with 1 L of TFSW for a 10 minutes sperm-egg contact 187 time to avoid polyspermy (Williams et al., 1997). Then, fertilized oocytes were removed and 188 washed twice with TFSW before being distributed (~  $10^5$  oocytes per container) in ten 189 different 1 L autoclaved glass containers filled with 500 mL of TFSW and placed in the dark 190 at 15°C. 191

192

193 Experimental design for larval rearing

194 After 48h post fertilization, TFSW was changed every two days with embryos retained and

195 washed in a 63-µm mesh. Some subsamples were fixed in 4% formaldehyde for further

observations. From day 4, the larvae began to secrete a lot of mucus, and to avoid clogging,they were gently resuspended with Pasteur pipette every day until day 12.

198 On day 6, the ten glass containers (1 L filled with 500 mL of TFSW) were placed in three

different thermostatically controlled rooms with respectively 3 glass containers at 13°C and

200  $17^{\circ}$ C, and 4 glass containers at  $15^{\circ}$ C.

After 24h of acclimation of these new temperature conditions at day 7, one glass container per room at 13°C and 17°C and two glass containers at 15°C were supplemented with a solution of microalgae (4.10<sup>4</sup> cell/mL concentration of RGcomplete APBreed<sup>TM</sup>, Planktovie ltd.) every two days and called the 'fed' conditions, while the remained containers at 13°C, 15°C and 17°C were called the 'non-fed' conditions. TFSW was changed initially every two days but after day 22 it was extended to three to five days. The experiments lasted for 50 days.

207

208 Monitoring of larval morphology and biometry

The larval development from artificially fertilized oocytes was monitored daily for the first 209 three weeks, then twice a week thereafter, using the optic microscope equipped with Motic 210 211 Image Plus© 3.0 software. Times required reaching the following stages of trochophore and metatrochophore were recorded for each temperature (13°C, 15°C and 17°C) and food 212 213 conditions ('fed' and 'non-fed'). For each temperature condition, fifteen to thirty larvae per glass container were collected at random and sacrificed for morphological observations and 214 biometry. The selected larvae were anesthetized (Gaudron & Bentley, 2002). The observation 215 216 of the number of setigers (segments bearing setae), as well as the opening of the mouth, the anus and the appearance of the digestive tract were carried out using the optic microscope. In 217 addition, taking photographs allowed to measure the total length of each larva (Motic Image 218 Plus© 3.0 software). 219

Scanning electron microscope (SEM) was used for better visualization of ontogeny. 220 221 For this, some larvae fixed in 4% formaldehyde were washed in MilliQ water (Millipore) in 40-µm mesh, and were gradually dehydrated by placing them successively for 1 hour in 222 ethanol (Merck, Normapur) baths ranging from 30% to 100% with a step of 10%. Following 223 this dehydration, and in order to fix and dry the larvae, they were put twice in a row, for one 224 hour, in a bath of hexamethyldisilane (HMDS, Molekula). The larvae were collected 225 226 individually using micro forceps and stuck on aluminium stubs (Agar Scientific) with double 227 sticky carbon tabs (Agar Scientific), which was finally sputter coated under Argon flow with Au/Pd (Polaron SC 7620) for 90 seconds. SEM observations were carried out under the SEM 228 229 LEO 438 VP using a secondary electron detector for topography at 20 keV.

230

231 Data analyses

232 Definition of 'birth'

The date of the first exogenous food intake, called 'birth' (DEB theory; Kooijman, 2010) in 233 our study, corresponds to the concomitant appearance of the opening of the mouth, of the anus 234 235 and the appearance of the gut. Initially Arenicola marina larvae are lecithotroph living on maternal reserve and this is called the 'embryo' stage in the DEB theory (Kooijman, 2010) 236 237 having a von Bertalanffy growth curve. Then when the larva starts to feed on exogenous food (planktotroph) by developing a functional gut, the growth is exponential until the end of the 238 metamorphosis. The transition between a lecithotrophic larva and the feeding larval stage has 239 240 been described for each temperature conditions through microscopic observation.

241

242 Biphasic bioenergetic modelling

Larval growth was modelled using a biphasic time-dependent model described by a set of twoequations. The change depends on the time of 'birth' (*tb*), where growth before 'birth' follows

- the laws of von Bertalanffy (von Bertalanffy, 1957) (Equation 1) and, after 'birth' it is
- exponential (Equation 2).
- 247 For length data, the growth equation is written as follows:

248 
$$L_1(t) = L_{inf} - (L_{inf} - L_0) * exp^{-bt}$$
 with  $L_{inf} = \frac{a}{b}$  for  $t \le tb$  i.e. before 'birth' (1)

249 
$$L_2(t) = L_1(tb) * exp^{ct}$$
 for  $t > tb$  i.e. after 'birth' (2)

250  $L_1(t)$  and  $L_2(t)$  are lengths as a function of time (*t*) with  $L_0$  and  $L_1(tb)$  are length at time 0 and 251 at *tb* respectively.  $L_{inf}$  is the asymptotic length, *a* and *b* are the size-specific rates of energy 252 acquisition and energy use for body maintenance (the von Bertalanffy growth rate between 253 fertilization and *tb*), respectively, and *c* the exponential growth rate after *tb*.

254

#### 255 Temperature range for metabolic responses in larval stages

All metabolic rates depend on body temperature (Kooijman, 2010), and in ectotherms it corresponds to the external temperature such as in polychaetes. Thus, a temperature correction (TC) is usually applied on metabolic rates using the Equation (3), where  $T_A$  is the Arrhenius temperature (in K),  $T_{ref}$ , the reference temperature (293.15 K), and *T* is the experimental temperature (in K):

261 
$$TC = \exp\left(\frac{T_A}{T_{ref}} - \frac{T_A}{T}\right)$$
(3)

Outside the lower and higher boundaries of the species-specific temperature tolerance range (respectively  $T_L$  and  $T_H$ ), the TC shape differs and is calculated adding an extra term to the Equation (3) as presented in Equation (4), with  $T_{AL}$  the Arrhenius temperature below the lower boundary of the species-specific temperature tolerance range (in K) and  $T_{AH}$  the Arrhenius temperature above the higher boundary of the species-specific temperature tolerance range (in K) (Kooijman, 2010).

$$269 \quad \exp\left(\frac{T_A}{T_{ref}} - \frac{T_A}{T_r}\right) \left[1 + \exp\left(\frac{T_{AL}}{T_{ref}} - \frac{T_{AL}}{T_L}\right) + \exp\left(\frac{T_{AH}}{T_H} - \frac{T_{AH}}{T_{ref}}\right) \right] (4)$$

A simpler version of this equation for the higher boundary of the temperature tolerance rangeonly is as follows:

273 
$$TC = \exp\left(\frac{T_A}{T_{ref}} - \frac{T_A}{T}\right) * \left[ \frac{1 + \exp\left(\frac{T_{AH}}{T_H} - \frac{T_{AH}}{T_{ref}}\right)}{1 + \exp\left(\frac{T_{AH}}{T_H} - \frac{T_{AH}}{T_{ref}}\right)} \right]$$
(5)

274 The Arrhenius temperature of A. marina has been previously estimated, using Equation 3, together with other DEB parameters using the DEBtool package (De Cubber et al., 2019; 275 Marques et al., 2018). In addition, the temperature tolerance range and the Arrhenius 276 temperatures of Equation 4 have been estimated for the species from data collected mainly in 277 juveniles and adults (De Cubber et al., 2020). Hence, new data on larval stages of A. marina 278 were used to re-estimate the Arrhenius temperatures i.e.  $T_A$ ,  $T_{AH}$  and to estimate the higher 279 280 boundary of the temperature tolerance range,  $T_H$ , using Equation 5. As no data were available below 5°C, it was not possible to estimate the lower boundary of Equation 4. The new data set 281 282 consisted in the parameters (a, b and c; Equations 1 and 2) of the biphasic growth model at 13°C, 15°C and 17°C as well as the data from several fertilization success experiments carried 283 out at 5°C, 10°C, 13°C, 15°C, 18°C, 20°C and 22°C by Lewis et al. (2002). Each data set was 284 285 standardized by its maximum value to get values between 0 and 1 in line with the temperature correction (Equation 5). 286

288 Statistics and fittings

289 All growth curve fitting processes and associated statistics were coded in R version 4.0.3 (2020). A nonlinear least squares method (package 'nls2'; Grothendieck, 2013) was used to fit 290 Equations 2 and 5 as it allows multiple starting values to avoid local minima problems in 291 parameter estimation. This package provides parameter best estimates and standard errors, 292 293 and parameter significances by *t*-test. Further, bioenergetics models were tested for either 294 differences in the temperature effect (3 modalities) or differences in the food condition ('fed' and 'non-fed') within each temperature (2 modalities) following the method of Ritz & 295 Streibig (2008) and using analysis of variance (ANOVA). For the temperature factor, the sum 296 of the residual sum of squares (RSS<sub>ind</sub>) of the three fitted models for each temperature (3 297 parameters per model, 9 in total, 'n\_par<sub>ind</sub>') were compared to the RSS<sub>all</sub> of a model grouping 298 all data and fitted with only 3 parameters ('n\_par<sub>all</sub>'). For the food condition factor, we 299 300 assumed there was no effect of food condition before tb and then, the von Bertalanffy phase of the biphasic model has L<sub>inf</sub> and b as common parameters for a given temperature. Hence, 301 the sum of RSS<sub>ind</sub> of the two fitted models for each food condition (2 common parameters 302 plus one c parameter per model, 'n\_par<sub>ind</sub>' = 4) were compared to the  $RSS_{all}$  of a model 303 grouping all data for a given temperature and fitted with only 3 parameters (' $n_{all}$ ' = 3). 304 305 The *F* statistic was calculated as follows:

$$F = \frac{\frac{RSS_{all} - RSS_{ind}}{(N - n_par_{all}) - (N - n_par_{ind})}}{\frac{RSS_{ind}}{N - n_par_{ind}}}$$

With *N* the total number of individuals. The *P* value was then determined by searching for the *F* value in the *F* distribution with degrees of freedom  $(n_par_{ind} - n_par_{all}, N - n_par_{ind})$ using the function 'pf' of the R statistical package.

310 Results

311 Effect of temperature on larval development of *Arenicola marina* 

According to the 13°C, 15°C and 17°C exposed temperature respectively, the chronology of 312 313 A. marina larval development (Fig. 1) and their biometry were recorded (Table 1). The fertilized oocytes had an average diameter of  $176 \pm 6 \,\mu\text{m}$  and develops to embryo by cell 314 division during embryogenesis (Fig. 1A). Larvae hatch at the trochophore stage (Fig. 1B) at 315 the end of the gastrulation at 3 days post fertilization (dpf) with a mean total length of  $169 \pm$ 316 317 14 µm (Table 1). The larvae developed their first setiger at 6 dpf (Fig. 1C) with a mean total length of  $255 \pm 28 \,\mu\text{m}$ , becoming a metatrochophore. All larvae were still conditioned at 318 15°C at that time (Table 1). After 6 dpf to 50 dpf, the larvae were raised to three different 319 temperature conditions (13°C, 15°C and 17°C), and the appearance of new setigers (up to 10 320 setigers) were not tuned between the three treatments. Indeed, at 15°C the larvae have 321 322 developed 4 setigers (S) at 14 dpf and 6 S at 24 dpf (Table 1; Figs. 1F&H). While the larvae placed at 13°C needed 16 and 24 dpf to reach 4 S and 6 S respectively (Table 1), those placed 323 324 at 17°C reached 4 S and 6 S at 14 and 21 dpf respectively (Table 1). At 50 dpf, larvae had 325 reached 7 S (Fig. 1I) with a mean total length of  $780 \pm 130 \,\mu\text{m}$  (Table 1) at  $13^{\circ}\text{C}$ , 8 S (Fig. 1J) with a mean total length of  $746 \pm 258 \,\mu\text{m}$  (Table 1) at  $15^{\circ}\text{C}$  and  $10 \,\text{S}$  with a mean total 326 length of  $544 \pm 186 \,\mu\text{m}$  (Table 1) at  $17^{\circ}\text{C}$ . Although, there was a time lag of larval 327 development as a function of temperature, larvae had equivalent size for each stage. Indeed, 328 for example for the 4 S stage, larvae measured  $471 \pm 69 \ \mu\text{m}$ ,  $476 \pm 33 \ \mu\text{m}$  and  $454 \ \mu\text{m} \pm 31$ 329 µm at 13°C, 15°C and 17°C respectively (Table 1). 330 331

332







- 337 Metatrochophore with C) 1 setiger (segment with chaetae); D) 2 setigers; E) 3 setigers; F) 4 setigers;
- 338 G) 5 setigers; H) 6 setigers; I) 7 setigers; J) 8 setigers.

- 340 Effect of diet on larval development of Arenicola marina
- 341 The dietary transition between lecithotrophic larvae to planktotrophic larvae ('birth') occurred
- at a size between ~ 450–500  $\mu$ m, regardless of temperature (Fig. 1F; Table 1; 1<sup>st</sup> time that a

row has 'yes' in the last two columns). In terms of duration, the age at 'birth' has taken place at 17 dpf for the larvae reared at both 15°C (4 S stage) and 17°C (5 S stage) and, at 21 dpf for those placed at 13°C (5 S stage) (Table 1).

Growth retardation was observed visually at 50 dpf between larvae fed with 346 microalgae ('fed') and those non-feds whatever the temperature treatments (Fig.2). For the 347 three-temperature conditions, the mean total lengths of larvae at 50 dpf of the 'non-fed' 348 conditions were lower than those of the 'fed' conditions (Table 2), but at 13°C mean total 349 length ( $659 \pm 96 \mu m$ ) of the 'non-fed' condition, was not significantly different than that of 350 the 'fed' condition ( $800 \pm 129 \ \mu m$ ) (*t*-test; P = 0.08); at 15°C, the mean total length of larvae 351 352 from 'non-fed' condition (487  $\pm$  110  $\mu$ m) was highly significantly different than that of the 'fed' condition (746  $\pm$  258 µm) (*t*-test; *P* < 0.001); at 17°C, the mean total length of larvae 353 from 'non-fed' condition (506  $\pm$  212 µm) was not significantly different than that of the 'fed' 354 355 condition (544  $\pm$  186 µm) (*t*-test; P = 0.56). It seems that some shrinkage had occurred in larvae from the 'non-fed' treatments between 43 dpf to 50 dpf both at 15°C and 17°C (Table 356 2). 357



Figure 2. Images obtained with an optic microscope of *Arenicola marina* larvae at different food levels and temperature conditions at 50 days post-fertilization. A) 13°C and 'fed' conditions; B) 15°C and 'fed' conditions; C) 17°C and 'fed' conditions; D) 13°C and 'non-fed' conditions; E) 15°C and 'nonfed' conditions; F) 17°C and 'non-fed' conditions.

Effect of temperature and diet conditions on bioenergetic of Arenicola marina larvae 363 Temperature had a significant effect on the biphasic growth models ( $F_{(6, 1338)} = 9.72$ ; P < 100364 0.001). In the first phase of the model (von Bertalanffy), growth rate b gave better 365 performance at  $15^{\circ}$ C (0.263 d<sup>-1</sup>) and  $17^{\circ}$ C (0.216 d<sup>-1</sup>) compared to  $13^{\circ}$ C (0.107 d<sup>-1</sup>), whereas 366 in the second phase of the model, exponential growth rates c were greater at  $13^{\circ}$ C (0.012 d<sup>-1</sup>) 367 and  $15^{\circ}$ C (0.013 d<sup>-1</sup>) compared to  $17^{\circ}$ C (0.006 d<sup>-1</sup>) (Table 3; Fig. 3). The effect of food on the 368 biphasic growth model is highly significant at  $15^{\circ}$ C ( $F_{(1,556)} = 59.44$ ; P < 0.001), where the 369 growth model gave better results in 'fed' conditions compared to the 'non-fed' condition 370 (Fig.3B). The effect of food is marginally significant at  $13^{\circ}$ C ( $F_{(1, 285)} = 2.77$ ; P = 0.097), but 371 372 still the biphasic growth model gave better performance in 'fed' condition compared to the 'non-fed' condition (Fig.3A). At 17°C, there is no effect of the food conditions on the 373 biphasic growth model ( $F_{(1, 494)} = 0.0$ ; P = 0.98), where both biphasic models were similar 374 375 given bad performance regarding larval growth (Table 3; Fig.3C).

376

377 Temperature correction on metabolic rates of *Arenicola marina* across different life-history378 stages

The estimates of the biphasic larval growth models (Table 3) along with data from Lewis et 379 380 al. (2002) after being standardized by their maximum values, helped to re-estimate the temperature corrections using Equation 5. As  $T_{AH}$  was non-significant in the first regression 381 fit (P = 0.14),  $T_{AH}$  (82380 K) from De Cubber et al. (2020) was set in the Equation 5. New 382 temperature corrections were calculated with a new  $T_A$  equaled to 6661.79 K (± 1241.5; P <383 0.001) and a new  $T_H$  equaled to 294.44 K (± 0.42; P < 0.001). Overall, these new Arrhenius 384 temperature datasets were different from those of De Cubber et al. (2020), where the  $T_H$  from 385 this study (blue line; Fig.4) issued from larval metabolic rates, was lower than that of De 386 Cubber et al. (2020) issued from juvenile/adult metabolic rates (black line; Fig.4), and the  $T_A$ 387

from our datasets (slope of the blue line on the left part of the curve; Fig.4) was higher than
that of De Cubber et al. (2020) (slope of the dark line on the left part of the curve; Fig.4).







- 412 indicated by the red dotted vertical line (17 days); C) At 17°C, where the age at first food intake
- 413 ('birth') is indicated by the dotted red vertical line (17 days). Lines are simulations of the models:
- classic von Bertalanffy (first phase of the biphasic growth model) and exponential (second phase of
- the biphasic growth model). Larval growth for the 'fed' condition is in black, and for the 'non-fed'
- 416 condition is in blue.



#### 417

Figure 4. Temperature corrections in different life stages of *Arenicola marina*. The black line
represents the temperature correction used in the abj-DEB model mostly on juvenile/adult stages
from De Cubber *et al.* (2020) and the blue line, the temperature correction using datasets from
fertilization success rate (circle, Lewis *et al.*, 2002) and from this study on larval growth with
parameters from the first phase of the biphasic growth model (*a* = black triangle; *b* = black square;
equation (1) and from the second phase of the biphasic growth model (*c* = black diamond; equation
(2)) at several temperatures.

425

#### 426 **Discussion**

- 427 The present work shows that a cohort of Arenicola marina was successfully fertilized in vitro
- 428 getting the embryo stage and then reared for fifty days under favorable experimental growth
- 429 conditions allowing the cohort to hatch to different larval stages including both a
- 430 lecithotrophic stage using maternal reserves and then, using the exogenous food provided by
- the algal culture being then planktotrophic. This experiment enabled to strengthen knowledge
- 432 on the first life-history stages of the lugworm species *A. marina*, and in particular the precise

age and length of the occurrence of the 'birth' stage with the biphasic growth before and after
'birth' under temperature and food control conditions. These data will be useful to consolidate
the abj-DEB model developed by De Cubber *et al.* (2019; 2020).

436

437 Early larval stages of *Arenicola marina* and 'birth' stage

Most of earlier work on developmental larval stages of A. marina were reported (Newel, 438 439 1948, 1949; Farke & Berghuis, 1979a,b) at a time that the species delimitation between A. marina and A. defodiens, a sympatric species that might occur at some beach in European 440 441 marine habitats, was not yet known (Cadman & Nelson-Smith, 1993; De Cubber et al., 2018) 442 mixing the different ontogeny larval stages between the two species. The best study on larval development was carried out by Farke & Berghuis (1979b) in laboratory where authors 443 develop a genius microsystem enabling mature adult lugworms (supposedly A. marina) to 444 spawn and larvae to develop in the laboratory. However, timing of spawning events, larvae 445 occurrence and control of temperature conditions could not be recorded precisely. Despite 446 447 this, previous authors (Farke & Berghuis, 1979b) described nicely the behavior, habitats and biometry of three larval stages of A. marina. Newly hatched trochophore larvae were in 448 female gallery and had a size around 0.25 mm in length. In our study the trochophore larval 449 450 stage was lesser in length and it was the larvae of one setiger that reached 0.25 mm. Metatrochophores with 3 setigers were seen swimming by ciliary movements and measured 451 around 0.5 mm (Farke & Berghuis, 1979b). In our study the 3S larval stage was ~ 0.4 mm 452 closed to what was measured by Newel (1948, 1949) from its in situ sampling larvae of A. 453 454 marina. After this 3S stage, larvae started to secrete a mucus tube in order to adhere to hard 455 substrate and they changed their behavior (Farke & Berghuis, 1979b). Larvae could leave their mucus tube in order to crawl and feed on particles deposited around the tube being 456 457 deposit-feeder (Farke & Berghuis, 1979b). Only larvae with 6 setigers were shown to contain

food particles in their gut with a size of 0.8 mm (Farke & Berghuis, 1979b). In our study the first food intake ('the birth stage') was observed earlier at the 4S/5S larval stage at a size between 450–500  $\mu$ m. Marty et al. (1997) had followed the appearance of setigers following time in the larvae of the polychaete *Hediste diversicolor*. Larvae of 3S (425 ± 30  $\mu$ m) were starting to feed ('birth') on non-fertilized oocytes in females gallery being cannibalistic and adelphophagic. This length at first feeding is very close to that observed in the larvae of *A*. *marina*.

Within an abj-DEB model, two primary parameters depends on the metabolic 465 acceleration  $(s_M)$  that occurs between the 'birth' stage to the end of the metamorphosis 466 (Kooijman, 2010; 2014): 1) The maximum assimilation rate after metamorphosis  $\{\dot{p}_{Am}\}_i =$ 467  $\{\dot{p}_{Am}\}_b s_M$  and; 2) The energy conductance values  $(\dot{v})$  after metamorphosis  $\dot{v}_j = \dot{v}_b s_M$ . The 468 metabolic acceleration is calculated as the ratio of the structural length at metamorphosis to 469 the structural length at 'birth':  $s_M = L_j/L_b$ . Within the abj-DEB model developed on A. 470 471 marina (De Cubber et al., 2019), the physical length at 'birth' ( $Lw_b$ ) was set at 230  $\mu$ m (twice lower to what is observed in this study) and this might have changed the estimation of the 472 metabolic acceleration  $s_M$ . In this study, we managed to describe precisely the length at 473 'birth' (~450 μm) and this will complete the dataset of the abj-DEB model of A. marina 474 developed by De Cubber et al. (2019). 475

476

477 Effect of abiotic factors on the first food intake ('birth') in Arenicola marina

When the larvae hatch at the trochophore stage, and until the development of the complete digestive tract occurring at the 4S/5S metatrochophore stage, the larvae draw their energy from the yolk reserves (lecithotrophy) for growth, maintenance and the complexity of its maturity in DEB theory (Kooijman, 2010). Thus, the availability of food in the environment has no influence on the transition from the lecithotrophic stage to the planktotrophic stage

('birth'), but temperature does. According to our results, 'birth' appeared earlier in 483 metatrochophores subjected to warmer temperatures (17 days at both 15°C and 17°C) 484 compared to lower temperature (21 days at 13°C). This is not in line with the age at 'birth' 485 486 estimated by the abj-DEB model proposed by De Cubber *et al.* (2019), where simulation carried at 10°C gave a 'first feeding' at 10.52 days post fertilization, twice much lower than 487 that observed at 13°C. However, when De Cubber et al. (2019) simulated the temperature 488 conditions for a whole year at Wimereux (Eastern English Channel) using real in situ data 489 from 5.5°C to 20°C, a closer simulated value of the age at 'birth' was estimated (15.5 days 490 closed to the 17 days observed for 15°C in our experimental set up). In the field, Newell 491 492 (1948, 1949) observed metatrochophores of Arenicola sp. ready to become planktotrophic at 2-3 weeks post spawning at Whistable (UK). In this study we managed to describe precisely 493 494 the age at 'birth' for three different temperatures and this will complete again the dataset of 495 the abj-DEB model of A. marina developed by De Cubber et al. (2019).

496

497 Abiotic factors on growth rates of *Arenicola marina* larvae

The increase in seawater temperature has induced an acceleration of larval 498 development giving at 50 days post fertilization, metatrochophores with more developed 499 500 setigers (10 S) in higher temperature conditions (17°C) compared to lower temperature; e.g. at 13°C only metatrochophores with 7 segments with chaetae were recovered. Thus, larvae 501 reared at 17°C changed larval stages faster than those exposed at 13°C meaning the energy 502 503 allocated to the complexity of the larvae was greater ( $E_H$  in DEB theory; Kooijman, 2010). However, the mean total length of the larvae reared at 17°C (~ 544 µm) was lower compared 504 505 to those reared at 13°C (~ 800 µm) at 50 dpf meaning in DEB interpretation that less energy was allocated to somatic growth while more energy was allocated to the complexity of the 506 507 larvae reared in higher temperature. The discrepancy in mean length was enhanced by the

poor food conditions treatment ('non-fed') that induced a kind of starvation at 50 dpf for both 15°C and 17°C treatments. In DEB theory (Kooijman, 2010), energy is needed in priority for maintenance of maturity and growth when less energy is available from mobilization; what is seen here is the larvae seem to shrink and some lysis of cells might have occurred.

In this study, the first phase (von Bertalanffy growth) of the biphasic growth model of 512 513 A. marina larvae that encompasses trochophores and metatrochophores up to 3S (before 'birth') was better at 15°C and 17°C. These larval stages occur first within the female gallery 514 515 on the intertidal foreshore and then disperse in the water column. Then, after 'birth', at the larval stage of 4S, the second phase (exponential growth) of the biphasic growth model was 516 517 greater at 13°C and 15°C, where at these larval stages, A. marina larvae are living on the subtidal areas. For both biphasic growth phases, the optimal temperature was shown to be at 518 15°C before and after 'birth'. Lewis et al. (2002) found for different populations of lugworms 519 520 in the UK that the optimal temperature for fertilization success (embryos stages) was between 15–18°C. Lewis et al. (2002) were quite astonished by their results as spawning periods of A. 521 522 marina occurred at lower temperature in the UK (10-12°C), where embryos develop in 523 female gallery on the intertidal habitat. Lewis et al. (2002) concluded that lugworms were not breeding at their optimal temperature and other selective pressures were certainly be acting. In 524 525 our study, the optimal temperature was found at 15°C and this, for others life-history stages (trochophores and metatrochophores) of A. marina that live not anymore on the intertidal area 526 but on the subtidal area (Farke & Berghuis 1979a; Newell 1948, 1949). At Wimereux 527 528 (Eastern English Channel), A. marina population spawns from the end of September to early October (De Cubber et al., 2018) where temperature drop from 15°C to 14°C but larvae 529 seems to be in their optimal temperature at least during the onset of larval development as 530 531 temperature fall in winter to temperatures up to 5.5°C (De Cubber et al., 2019). At a regional scale, other populations of A. marina breed later until mid-November on the Eastern English 532

Channel (De Cubber *et al.*, 2018). In mid-November, temperature is around 10°C as seen in the UK in Lewis *et al.* (2002). *A. marina* populations are widespread in Europe and some population live in South of Europe such as in Portugal (Pires *et al.*, 2015) where the mean seawater temperature is much higher in winter but in the range of the optimal temperature for larvae and in spring and summer in the range of juvenile/adult optimal temperature. This may explain the well-establishment of this species in South of Europe, where in Portugal the lugworm is seen as an invasive species (Pires *et al.*, 2015).

540

541 Applications in DEB theory and in aquaculture

Intertidal species (mostly ectotherms) such as polychaetes, bivalves and gastropods can 542 experience during low tide a great variation (up to 20°C) of daily temperature either in winter 543 or in summer (Seuront et al., 2019; Moisez et al., 2020; De Cubber et al., 2020) compared to 544 species living in a more stable daily temperature environment such as in the subtidal area. As 545 546 reported by Kooijman (2010), these species have enzymes involved in metabolic reaction that 547 function in this broad temperature range with the consequence to have a relatively low Arrhenius temperature  $(T_A)$  (around 6000 K), compared to species that live in more constant 548 daily temperature having a higher Arrhenius temperature (around 12 000 K). T<sub>A</sub> calculated 549 550 using DEB tool (Add-my-pet-database) of the polychaete Hediste diversicolor and the cockle bivalve Cerastoderma edule, living both on intertidal mud flat, were found respectively to be 551 4877 K and 5290 K respectively. De Cubber et al. (2020) have estimated a T<sub>A</sub> of 4014 K for 552 Arenicola marina, a correct value for an intertidal species. The calculation was based on 553 metabolic rates of life-history stages of the lugworms (embryos, juveniles and adults) that live 554 on the foreshore. In our study, a new set of Arrhenius temperatures ( $T_A$  and  $T_H$ ) was calculated 555 based on temperature corrections of metabolic rates of only early-life stages of A. marina 556

(embryos and larvae).  $T_A$  of early-life stages of A. marina, that spend most of their time in the

558	subtidal area (a more stable environment), as expected, was found higher (~ 6661 K)
559	compared to the $T_A$ (~ 4014 K) (De Cubber et al. 2020) of life-stages of A. marina that live on
560	the foreshore (a highly variable environment). Likewise, the higher boundary temperature
561	value ( $T_H = 294.4 \text{ k}$ ; ~ 21.25°C) of the early-life stages was lower than that of the
562	juvenile/adult stages ( $T_H = 297.7 \text{ k}$ ; ~ 24.55°C; De Cubber <i>et al.</i> , 2020). As already reported
563	by Kooijman (2010), larvae of intertidal species that live in pelagic environment, have a
564	higher Arrhenius temperature as this $T_A$ can change with the life stage of a species. We
565	therefore support the idea that two sets of Arrhenius temperatures should be used in all
566	intertidal Lophotrochozoan species that have a larval life in pelagic area when using an abj-
567	DEB model. As the authors are aware only one Arrhenius temperature is usually including
568	into any DEB model even if, a species may experience different temperature ranges during
569	their life cycle. For instance, in the mollusc bivalve Magdallena gigas, that is a commercial
570	species and intertidal, in the AMP database, the value of $T_A$ is set at 8000 K despite that Rico-
571	Villa et al. (2010) calculated a higher value of $T_A$ (11 000 K) for the larvae after rearing them
572	at 5 different temperatures from 17°C to 32°C.
573	A. marina has been cultured since the late 90' in Northeast England (Northumbland
574	Seabait ltd.) with a number of patents issued from this bait farming (e.g. Olive et al., 2001

Seabait ltd.) with a number of patents issued from this bait farming (e.g. Olive et al., 2001 (WO2003007701A2); Craig & Olive, 2005 (WO2005043994A1)). The initial purpose of the culture of lugworms in the UK was to support the demands of fishermen that were digging intensively the worms used for bait (Olive, 1993; 1994; Olive & Cowin, 1994). Recently, *A. marina* is reared in a farm in Noirmoutiers Island in West of France (Hemarina Ltd<sup>TM</sup>) for medical purposes, where a numerous of exiting research is carried out on the medical potential and application of the lugworm haemoglobin (Asong-Fontem *et al.*, 2021; Batool *et al.*, 2021; Le Daré *et al.*, 2021; Le Meur *et al.*, 2021). Our study on larval physiology

highlight that the optimal temperature for growth is around 15°C with a maximal tolerance of
21°C and this could have interesting application in aquaculture.

584

#### 585 Conclusion

Overall, our data on the early larval stages of A. marina will be valuable in improving the 586 existing abj-DEB model for this engineer species. These include life traits such as age at birth, 587 588 size at birth but also Arrhenius temperatures and length over time for two food levels. DEB modeling allows to predict functional traits of the species such as size at first maturity, life 589 span, number of oocytes during the whole life cycle (total reproductive output), growth rate, 590 591 maximum length (Lmax), etc. This model outputs can help marine conservation managers make decisions to preserve the A. marina population exploited by bait fishing. In particular, it 592 593 helps stakeholders to establish regulatory measures such as catch size limits or the number of individuals that can be harvested. One of the solutions to overexploitation of lugworms is 594 595 aquaculture farming. Our data underline that the optimal temperature for rearing lugworm 596 larvae is 15°C and that it is necessary to feed them with microalgae after the 'birth' period 597 which occurs 17 days after fertilization.

598

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603

604 Conflicts of interest

The authors declare no financial and personal conflict of interest.

606

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613	sets of Arrhenius temperatures for species living in different marine habitats during their life
614	cycle.
615	
616	Data availability statements
617	The data underlying this article are available in the article.
618	
619	References
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**Table 1.** Larval development of *Arenicola marina* at 13°C, 15 °C and 17°C. At 7 days post-fertilization

807 (Time), larvae were fed with microalgae. Larval stages correspond to the number of setigers (S).

808 Total length is the mean of the n replicates with its standard deviation (±).

Temperature	Time (days)	Larval stage	Total length	n	Mouth & anus	Presence of a
			(μm)		opening	digestive tract
15°C	0 – 2	Embryo	159 ± 8	73	No	No
	3	Trochophore	169 ± 14	91	No	No
	6	1S	255 ± 28	48	No	No
	9	2S	356 ± 15	8	No	No
	12	3S	383 ± 34	12	No	No
	14	4S	476 ± 33	23	Yes	No
	17	4S	465 ± 52	22	Yes	Yes
	20	5S	506 ± 66	16	Yes	Yes
	24	6S	499 ± 86	16	Yes	Yes
	43	7S	541 ± 157	6	Yes	Yes
	50	8S	746 ± 258	22	Yes	Yes
13°C	8	25	246 ± 11	5	No	No
	13	3S	415 ± 30	10	No	No
	16	4S	471 ± 69	8	Yes	No
	21	5S	510 ± 46	5	Yes	Yes
	24	6S	461	1	Yes	Yes
	43	7S	674 ± 76	4	Yes	Yes
	50	7S	800 ± 129	5	Yes	Yes
17°C	8	25	276 ± 19	11	No	No
	11	3S	371 ± 21	9	No	No
	14	4S	454 ± 30	19	Yes	No
	16	5S	445 ± 40	8	Yes	No
	17	5S	459 ± 45	10	Yes	Yes
	21	6S	588 ± 95	6	Yes	Yes
	37	7S	551 ± 155	18	Yes	Yes
	43	8S	586 ± 204	20	Yes	Yes
	50	10S	544 ± 186	21	Yes	Yes

**Table 2.** Larval development after 'birth' according to diet condition for each temperature treatment.

- 819 Times correspond to days post-fertilization. Total length is the mean of the n replicates with its
- 820 standard deviation (±). na for not available.

Temperature	Time (days	Total length (μm) for	Total length (μm) for
	post-	'non-fed' condition	'fed' condition
	fertilization)		
13°C	21	551 ± 74 ( <i>n</i> = 12)	505 ± 51 ( <i>n</i> = 4)
	24	543 ± 79 ( <i>n</i> = 13)	461 ± na
	27	594 ± 65 ( <i>n</i> = 8)	497 ± 64 ( <i>n</i> = 9)
	29	626 ± 132 (n = 7)	463 ± 25 (n = 2)
	37	651 ± 120 ( <i>n</i> = 14)	561 ± 64 ( <i>n</i> = 5)
	43	626 ± 102 ( <i>n</i> = 11)	674 ± 76 ( <i>n</i> = 4)
	50	659 ± 96 ( <i>n</i> = 12)	800 ± 129 ( <i>n</i> = 5)
15°C	17	465 ± 84 (n = 12)	465 ± 52 (n = 22)
	21	468 ± 80 ( <i>n</i> = 7)	472 ± 34 (n = 7)
	24	465 ± 71 ( <i>n</i> = 12)	499 ± 86 ( <i>n</i> = 16)
	27	511 ± 95 ( <i>n</i> = 19)	485 ± 78 (n = 13)
	29	514 ± 113 ( <i>n</i> = 15)	467 ± 68 (n = 15)
	37	556 ± 77 ( <i>n</i> = 13)	583 ± 85 ( <i>n</i> = 14)
	43	613 ± 154 ( <i>n</i> = 9)	541 ± 157 ( <i>n</i> = 6)
	50	487 ± 110 ( <i>n</i> = 23)	746 ± 258 (n = 22)
17°C	17	476 ± 57 ( <i>n</i> = 11)	459 ± 45 (n = 10)
	21	500 ± 70 ( <i>n</i> = 11)	588 ± 95 ( <i>n</i> = 6)
	24	420 ± 70 ( <i>n</i> = 10)	485 ± 66 ( <i>n</i> = 8)
	27	466 ± 133 ( <i>n</i> = 10)	505 ± 107 ( <i>n</i> = 12)
	29	463 ± 72 ( <i>n</i> = 11)	473 ± 57 (n = 8)
	37	514 ± 125 (n = 13)	551 ± 155 ( <i>n</i> = 18)
	43	687 ± 265 ( <i>n</i> = 16)	586 ± 204 ( <i>n</i> = 20)
	50	506 ± 212 ( <i>n</i> = 17)	544 ± 186 ( <i>n</i> = 21)

**Table 3.** Biphasic growth modelling parameters where *a* and *b* are from Von Bertalanffy's phase and

*c* results from the exponential phase. The parameter *a* was estimated from *L<sub>inf</sub>* and *b* using Equation
2.

	Treatment	Parameters	Unit	Value	Standard error	P-value
	13°C	L <sub>inf</sub>	μm	595.83	43.77	< 0.001
		a	µm/d	63.71	-	-
		b	/d	0.107	0.026	< 0.001
	Fed	С	/d	0.012	0.001	< 0.001
	Non-fed	С	/d	0.010	0.001	< 0.001
	15°C	L <sub>inf</sub>	μm	481.59	15.02	< 0.001
		а	µm/d	126.92	-	-
		b	/d	0.263	0.051	< 0.001
	Fed	С	/d	0.013	0.001	< 0.001
	Non-fed	С	/d	0.005	0.001	< 0.001
	17°C	L <sub>inf</sub>	μm	512.61	29.15	< 0.001
		а	μm/d	110.61	-	-
		b	/d	0.216	0.060	< 0.001
	Fed	С	/d	0.006	0.001	< 0.001
-	Non-fed	С	/d	0.006	0.001	< 0.001
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841 Figures caption

Figure 1. Scanning electron microscopy photographs of ten larval stages of Arenicola 842 marina. A) Embryo at early stage of cell division before hatching; B) Trochophore stage after 843 844 hatching; Metatrochophore with C) 1 setiger (segment with chaetae); D) 2 setigers; E) 3 setigers; F) 4 setigers; G) 5 setigers; H) 6 setigers; I) 7 setigers; J) 8 setigers. 845 Figure 2. Images obtained with an optic microscope of Arenicola marina larvae at different 846 food levels and temperature conditions at 50 days post-fertilization. A) 13°C and 'fed' 847 conditions; B) 15°C and 'fed' conditions; C) 17°C and 'fed' conditions; D) 13°C and 'non-848 849 fed' conditions; E) 15°C and 'non-fed' conditions; F) 17°C and 'non-fed' conditions. Figure 3. Evolution of the total length of Arenicola marina larvae at three different 850 temperatures following days post-fertilization. A) At 13°C, where the age at first food intake 851 852 ('birth') is indicated by the red dotted vertical line (21 days); B) At 15°C, where the age at first food intake ('birth'), is indicated by the red dotted vertical line (17 days); C) At 17°C, 853 where the age at first food intake ('birth') is indicated by the dotted red vertical line (17 days). 854 Lines are simulations of the models: classic von Bertalanffy (first phase of the biphasic 855 growth model) and exponential (second phase of the biphasic growth model). Larval growth 856 for the 'fed' condition is in black, and for the 'non-fed' condition is in blue. 857 Figure 4. Temperature corrections in different life stages of Arenicola marina. The black line 858 represents the temperature correction used in the abj-DEB model mostly on juvenile/adult 859 stages from De Cubber et al. (2020) and the blue line, the temperature correction using 860 datasets from fertilization success rate (circle, Lewis et al., 2002) and from this study on 861 862 larval growth with parameters from the first phase of the biphasic growth model (a = black triangle; b = black square; equation (1)) and from the second phase of the biphasic growth 863 864 model (c = black diamond; equation (2)) at several temperatures.

865 Tables caption	
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**Table 1.** Larval development of *Arenicola marina* at 13°C, 15 °C and 17°C. At 7 days post-

867 fertilization (Time), larvae were fed with microalgae. Larval stages correspond to the number

of setigers (S). Total length is the mean of the n replicates with its standard deviation  $(\pm)$ .

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870	<b>Table 2.</b> Larval development after 'birth' according to diet condition for each temperature
871	treatment. Times correspond to days post-fertilization. Total length is the mean of the n
872	replicates with its standard deviation $(\pm)$ . na for not available.

873

**Table 3.** Biphasic growth modelling parameters where *a* and *b* are from Von Bertalanffy's phase and *c* results from the exponential phase. The parameter *a* was estimated from  $L_{inf}$  and b using Equation 2.

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