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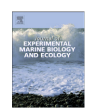
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## Oxygen and capacity limited thermal tolerance of the lugworm Arenicola marina: A seasonal comparison

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#### ABSTRACT

Lugworms Arenicola marina were collected from Arcachon Bay in two summers and winters of consecutive 22 years. The worms were acclimated to different temperatures (5 and 10 °C for winter animals and 15 °C for 23 summer animals). Each group was investigated over an experimental temperature range concerning its op- 24 timum in exercise performance, acute growth rate as well as respiration and ventilation activities to reveal 25 seasonal acclimatisation effects, potential inter-annual differences and the influence of laboratory acclima- 26 tion temperatures on the parameters of interest. The groups investigated at the two consecutive summers 27 yielded nearly identical results for ventilation and respiration activities. A clear seasonal difference developed 28 in exercise performance, with an optimum at lower temperatures in winter than in summer, irrespective of 29 acclimation temperature. Respiration and ventilation activities showed no significant differences between 30 winter specimens acclimated to 10 °C and summer specimens acclimated to 15 °C. However, an acclimation 31 temperature of 5 °C for winter animals caused noticeable differences to those acclimated at 10 °C. Acute 32 growth rates differed seasonally as well as between acclimation temperatures with the highest rates found 33 around 10 °C in summer and around 15 °C in winter. The lowest rates were recorded in winter worms accli- 34 mated to 5 °C. These acute patterns may reflect high thermal limits in warm acclimated winter worms and 35 temperature dependent shifts in energy demand in summer animals.

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## 1. Introduction

Arenicola marina is one of the most eminent secondary producers in the intertidal habitat. It is important for other epibenthic and infaunal animals aerating the habitat by sediment dwelling. The fresh weight of individuals may rise up to more than 30 g. Field studies in northern France showed that up to 346.6 kg lugworm faeces were produced per m<sup>2</sup> and year, which corresponds to a sediment layer of 21.5 cm height (Pollack, 1979). A. marina feeds on ciliates, microalgae and bacteria in the sediment and the overlying water. Bacterial biomass and chlorophyll a concentration in the sediment show a clear seasonal variability with a maximum occurring between April and October. In contrast, coastal surface waters show the highest bacterial biomass and concentrations of chlorophyll a in March to July (Hubas et al., 2007). Ciliates and other mesopsammon-organisms also show the highest density in summer (Pollack, 1979). The main food uptake by A. marina occurs during high tide, when the sediment is covered by surface water. A. marina not only exploits the nutrients enclosed in the upper sand layer by consuming the sand caving in from the surface like a funnel, but also extracts those from the surface water (Pollack, 1979) by generating a headward directed water current through its

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burrow (Wells, 1945). Suspended substances and planctonic organisms 62 are trapped in the sand of the burrow headshaft, which acts as a filter 63 (Krüger, 1957). The worm emits mucus to fill the interstices so that 64 small colloidal particles are also retained. This way, the food region is 65 enriched of organic material, which the lugworm ingests together 66 with the sand.

A. marina is very abundant in Arcachon Bay. It is found at any 68 beach (Boisseau, 1962) in densities up to 20 or sometimes 30 to 40 in- 69 dividuals per m<sup>2</sup>, mostly in the intertidal zone and around the island 70 Ile aux Oiseaux even in direct neighbourhood of oyster and eelgrass 71 beds (Amoureux, 1966). The studied population is located at La 72 Hume, a sheltered beach at the southern coast of Arcachon Bay. The 73 surface inhabited by lugworms is a band of approximately 300 m 74 width, bordered by a Zostera nana bed at the lower margin and a 75 salt marsh vegetation at the upper margin (Cazaux, 1966). The popu- 76 lation consists of at least three generations. The youngest generation 77 is found from the end of March onward in the highest zone of the in- 78 tertidal, close to the sandy beach. The older individuals with a body 79 weight of up to 6 g inhabit the lower intertidal (Cazaux, 1966). At 80 La Hume lugworms start the production of gametes around the end 81 of April, spawning occurs between the end of August and the begin- 82 ning of October. After reproduction, the oldest generation disappears 83 and the younger ones resume growing until spawning of the next 84 year (Cazaux, 1966). 85

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This strong seasonal pattern mirrors the temperature changes between seasons which are accompanied by a shift in the widths and positioning of thermal tolerance windows on a temperature scale. Seasonal changes in A. marina thermal tolerance have already been investigated in a North Sea population (Sommer, 2001; Sommer and Pörtner, 2004; Sommer et al., 1997; Wittmann, 2005; Wittmann et al., 2008). According to the concept of oxygen and capacity limited thermal tolerance, critical temperatures represent the threshold beyond which anaerobic metabolism is necessary for survival because the oxygen demand can no longer be met by aerobic processes (Frederich and Pörtner, 2000; Pörtner, 2001). A more or less parallel shift of both low and high critical temperature values was found with seasonal acclimatisation, characterised by anaerobic end product accumulation (Sommer et al., 1997). Investigations at the mitochondria level revealed that seasonal cold acclimatisation in North Sea lugworms involved a drop in mitochondrial density below summer values, combined with an increasing efficiency of aerobic energy production of each individual mitochondrion (Sommer, 2001; Sommer and Pörtner, 2004). Wittmann et al. (2008) compared thermal tolerance windows of North Sea lugworms during winter to those during summer and demonstrated a widening of the window, accompanied by a shift of critical temperatures towards higher values.

The present study was designed to assess the physiological responses accompanying seasonal acclimatisation and the biochemical changes reported above. Effects of temperature acclimation and inter-annual variation were investigated in a Southern population of the lugworm. Methods were chosen to investigate exercise performance capacity, metabolic energy demand and supply as well as somatic growth. Recordings of digging periods have already been established as a measure for muscular performance and were successfully applied to show differences in performance levels of lugworm populations from various latitudes (Schröer et al., 2009). The water volume which the lugworm pumps through its burrow gives some information about oxygen demand, as oxygen is extracted by gills and body surface from the bypassing water. Pumping frequency is changed to enhance or reduce the water volume flow through the burrow in order to adjust oxygen availability to the respective demand. The experimental setup simultaneously recorded water volume flow and oxygen content and also provided data for pumping frequency and oxygen extraction efficiency determination. Pumping frequency has already been used as a measure for performance depending on salinity (Shumway and Davenport, 1977) and temperature (Schröer et al., 2009; Wittmann, 2005; Wittmann et al., 2008). The mechanical aspects of the lugworm pump (Riisgård et al., 1996) and the biogeochemical consequences for the vented burrow fluids (Davey et al., 1990) have also been of interest. For the determination of acute growth optima, the incorporation of <sup>13</sup>C labelled phenylalanine was tracked by <sup>13</sup>C NMR spectroscopy as introduced by Wittmann et al. (2008).

## 2. Materials and methods

## 2.1. Animals

Specimens of the polychaete *A. marina* (L.) were collected in the intertidal zone at the sampling site in La Hume (44.65° N, 1.17° W) near Arcachon at the French Atlantic coast. Lugworms collected in August 2005 were used for investigations of respiration, ventilation and protein syntheses. For the same experiments during winter, animals were collected in January/February 2006. For yearly comparison the collection of animals was repeated in August 2006 and reinvestigated for potential annual changes. Digging performance was studied on animals from August 2006 and February 2007, for summer and winter, respectively. All worms were maintained in basins filled with natural sediment in a natural seawater flow-through

aquarium system at the Alfred Wegener Institute until experimental 149 use. The specimens collected in winter were divided between incubation temperatures of 5 °C and 10 °C, those collected in summer were 151 kept at 15 °C. All animals were exposed to a salinity of 32% and a 152 h/12 h light/dark cycle in the aquaria and fed with ground and 153 soaked Tetramin® flakes every other week.

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#### 2.2. Field measurements

In parallel to animal collection, biotic and abiotic field parameters 156 were recorded at the sampling site as described by Schröer et al. 157 (2009). In particular, temperatures in air, tidal puddles and sediment 158 were recorded with a thermometer (Testo 925, Testo, Lenzkirch, 159 Germany) using a special temperature-receiving element (6 mm 160 diameter, 500 mm length, TC Direct, Mönchengladbach, Germany). 161 Salinity was measured in tidal puddles by use of a multiple parameter 162 pocket measurement device (Multi 340i, WTW, Weilheim, Germany). 163 The length of the tail shaft of the worm's burrow was taken as a mea-164 sure of burrow depth by using a scaled metal stick inserted into the 165 opening of the burrow. For abundance recordings, a 1 m×1 m wooden 166 frame was placed onto the intertidal sediment at haphazard and the 167 number of faecal piles therein was counted. Bodyweight was measured using a scale (EMB 220-I, Kern, Balingen-Frommern, Deutschland).

## 2.3. Digging performance

Experimental temperatures were chosen at 4 °C steps, between -1 171 and 19 °C for the winter worms. Specimens sampled in summer were 172 measured at 7 to 27 °C. For initial short-term acclimation animals 173 were transferred to a plastic container placed into a temperature con-174 trolled aerated seawater bath. Temperature was changed at 2 °C h\_1^-1 175 starting from maintenance conditions and kept constant for at least 12 h at the new experimental temperature. After acclimation for at 177 least 12 h\_1 animals were transferred into the experimental setup 1 h 178 prior to measurements.

The experimental setup and procedure were the same as described 180 by Schröer et al. (2009). Briefly, the animals were positioned on the sed- 181 iment surface and the duration of each digging period was recorded 182 using a stopwatch. For an analysis of burrowing capacity during a 183 limited time window this routine was repeated for 90 min and the 184 number of digging periods was recorded. In total, five specimens of 185 each group were examined at each temperature (n=5). Summer 186 (15 °C acclimated) data were published in our previous work recently 187 (Schröer et al., 2009).

### 2.4. Respiration and ventilation experiment

Analyses of respiration and ventilatory activity were carried out as 190 described previously (Wittmann et al., 2008). Briefly, measurements 191 were performed in the dark using artificial burrows consisting of 192 straight Perspex tubes with a rough inner surface. As in their natural 193 burrows, animals generated a water current to provide themselves 194 with oxygen. Air saturation of incurrent and excurrent water was monitored continuously with oxygen micro-optodes (PreSens, Regensburg, 196 Germany). The volume flow produced by the worms was measured 197 using an electromagnetic flowmeter (inner diameter of probe head: 198 3 mm, RT-500, Hugo Sachs Elektronik, March-Hugstetten, Germany).

Experiments started at a temperature of 4.5 °C for winter animals 200 kept at 5 °C and at 10.7 °C for those kept at 10 °C. Summer animals 201 were tested beginning at 15 °C. Temperature was changed at a rate 202 of 1 °C h $^{-1}$  (first lowered) by steps of 3 °C and kept constant for 203 6 h. Lugworms collected in winter were exposed to a temperature 204 range from -0.0 to 22.8 °C, summer worms experienced a range 205 from 2.8 to 26.1 °C. Mean oxygen partial pressure of incurrent and 206 excurrent water ( $P_{\text{IO}_2}$  and  $P_{\text{EO}_2}$ , kPa) and weight specific volume flow 207 ( $V_{\text{w}}$ , ml  $h^{-1}$  g $^{-1}$ ) were calculated for the last 3 h of each incubation 208

period. From these data oxygen consumption  $(M_{O_2}, \mu mol\ O_2\ h_{\perp}^{-1}\ g_{\perp}^{-1})$  and the extraction coefficient (%) were determined. The analysis also included the mean volume flow during the phases of active ventilation (active volume flow,  $\min_{\perp}^{-1}$ ) and the number of contraction waves of the body wall (pumping frequency,  $\min_{\perp}^{-1}$ ). The water volume transported per peristaltic wave of the body wall musculature (wave volume, ml, see Wittmann et al., 2008) was calculated from volume flow recordings during active ventilation periods. Results from summer 2006 animals were published before (Schröer et al., 2009).

#### 2.5. Protein biosynthesis

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Analyses of protein biosynthesis were carried out as described previously (Wittmann et al., 2008) and following the principles outlined by Langenbuch et al. (2006). Briefly, uniformly labelled  $^{13}\text{C-L-phenylalanine}$ , dissolved in filtered seawater (75 mmol  $\text{L}^{-1}$ ), was injected at 40  $\mu$  g\_ $^{-1}$  body weight into the coelomic cavity of the lugworms. The animals were then inserted into their artificial burrows and incubated at temperatures of -0.9, 3.5, 6.9, 11, 15.2 and 19.2 °C for winter animals and 7.4, 10.2, 14.3, 17.8 and 22.8 °C for summer animals while they were actively ventilating their burrows. For animals collected in summer the experiment was carried out with 8 animals at each temperature. Animals were exposed to incubation temperatures for 24 h, then injected and incubated for another 30, 60, 90, 120, 180, 240, 300 and 360 min\_ respectively. For winter animals, 12 specimens were used at each temperature and 2 of them pooled after 30, 60, 120, 180, 240 and 360 min\_ respectively.

After the respective incubation time, the cuticulo-muscular tube was frozen in liquid nitrogen. The frozen tissue was ground under liquid nitrogen and extracted with TCA (trichloroacetic acid). The homogenate was centrifuged and supernatant and pellet were treated differently. The supernatant representing the cytosolic fraction and containing low molecular weight constituents was neutralised, dried and dissolved in D<sub>2</sub>O (deuterium oxide).

The pellet was washed, suspended in distilled water and neutralised. After centrifugation, the supernatant containing water-soluble proteins was removed and stored while the pellet was boiled in a water bath with 1 M NaOH. The water-insoluble proteins were then added to the water-soluble protein fraction and both were dried. The total protein was dissolved in D<sub>2</sub>O.

Both cytosolic and protein extracts were measured in a NMR Spectrometer (9.4 T Avance, Bruker, Rheinstetten, Germany) at frequencies of 100.6 MHz for <sup>13</sup>C spectra. The same parameters of the NMR recordings were used as described in Wittmann et al. (2008).

The protein contents of the protein extracts were quantified according to Bradford (1976). The amount of incorporated <sup>13</sup>C-L-phenylalanine was obtained as described by Wittmann et al. (2008). The calculated amount of incorporated <sup>13</sup>C-L-phenylalanine was related to the respective protein content of the sample and plotted against incubation temperature for incubation time. Due to the insensitivity of the NMR technique and the fact that only one sample per individuum could be analysed we could not track the time-dependent increase of incorporated <sup>13</sup>C-L-phenylalanine in sufficiently high resolution. Inter-individual differences exceeded those determined over time, which is why data points were pooled into two time frames: short incubation period (30 to 120 min) and long incubation period (180 to 360 min) (see also Wittmann et al., 2008).

#### 2.6. Statistics

Statistical analysis was performed using GraphPad Prism version 4.0c for Macintosh (GraphPad Software, San Diego, California, USA). Nonlinear regression curves were fitted to temperature dependent burrowing activities, using the equation  $EP(T) = F_1(T) + F_2(T) = F_1(T) + F_2(T)$ 

 $(A_1e^{B1T}+C_1)+(A_2e^{B2T}+C_2)$  according to Pörtner and Knust (2007). 271 For weight specific volume flow data the equation  $V_w(T) = Ae^{BT} + C$  272 was used as described in Wittmann et al. (2008). Significant devia- 273 tions of V<sub>w</sub> values from the hypothetical value extrapolated from 274 the exponential relationship were identified by one-sample Student's 275 t-test. One-way ANOVA was performed to detect significant changes 276 in pumping frequency, extraction coefficient and protein synthesis 277 values over the temperature range within each group (F-test). Two- 278 way ANOVA combined with a Bonferroni posthoc test was applied 279 to detect significant differences in pumping frequency between the 280 groups at each temperature.  $Q_{10}$  values were calculated for specific 281 temperature ranges using standard procedures as in Sommer and 282 Pörtner (2002). One-way ANOVA and Tukey's posthoc test were 283 used to identify differences in field data between the groups. Statis- 284 tical significance was identified at the p $\leq$ 0.05 level. All data are 285 given as means  $\pm$  SE if not stated otherwise.

## 3. Results

#### 3.1. Field measurements

Biotic and abiotic field parameters (Table 1) were recorded to 289 characterise the natural habitat conditions of the population at the 290 time of collection during the respective season. Salinity was signifi- 291 cantly (p<0.001) lower in winter than in summer, whereas vari- 292 ability between the two consecutive years was small. Temperatures 293 in air, tidal puddles and sediment (20 cm depth) differed significantly 294 (p<0.001) between winter and summer. Differences between the two 295 winters or the two summers, respectively, were much smaller, although 296 some significant deviations could still be found (air temperature in 297 winter 2006 vs. winter 2007: p<0.001; temperature in tidal puddles 298 summer 2005 vs. summer 2006: p<0.001; sediment temperature in 299 20 cm depth: p<0.001 between all groups), as data were only available for the week of animal collection. Burrow depth (length of the 301 tail shaft) did not show any seasonal differences. Abundance also 302 did not vary between winter and summer. A seasonal variation in 303 animal weight could be detected; specimens collected in summer 304 2006 were significantly heavier than animals collected in winter 305 (vs. winter 2007: p<0.001; vs. winter 2006: p<0.05). The value for 306 summer 2005 is not representative, as juveniles were comprised in 307 the measurement.

## 3.2. Digging performance

Fig. 1 shows the temperature dependent number of digging periods 310 performed during 90 minutes experimental intervals. Lugworms collected in winter and acclimated to 5 °C showed up to  $8.60\pm0.24$  digging periods  $90\,\mathrm{min}^{-1}$  at  $15.4\,^{\circ}\mathrm{C}$ . Digging performance decreased 313 rapidly towards higher temperatures and more slowly towards lower 314 temperatures. At -0.8, 3.0 and 7.0 °C, digging activity stayed on the 315 same level of about 4 periods  $90\,\mathrm{min}^{-1}$ . The regression analysis of 316 performance exhibited a maximum of 7.35 digging periods  $90\,\mathrm{min}^{-1}$  317 at  $14.2\,^{\circ}\mathrm{C}$ .

Winter animals acclimated to 10 °C showed similar digging per-  $^{319}$  formances. They displayed the highest number of digging periods  $^{320}$  (7.80  $\pm$  0.37 90 min $^{-1}$ ) at 15.3 °C. As seen in 5 °C acclimated worms,  $^{321}$  digging performance decreased rapidly towards higher temperatures  $^{322}$  and more slowly towards lower temperatures. In contrast to the latagraph terror values at  $^{-1.1}$ , 3.0 and 6.9 °C did not stay at the same level but  $^{324}$  increased steadily. The performance model indicated a maximum of  $^{325}$  7.48 periods 90 min $^{-1}$  at 13.3 °C.

Lugworms collected in summer and acclimated to  $15\,^{\circ}\text{C}$  per- 327 formed the highest number of digging periods  $(5.80 \pm 0.80\,90\,\text{min}^{-1})$  328 at  $23.2\,^{\circ}\text{C}$ , while the model predicted a maximum of 5.26 periods  $329\,90\,\text{min}^{-1}$  at  $19.6\,^{\circ}\text{C}$ . As observed in winter worms, performance capacity decreased more rapidly towards higher than towards lower  $331\,^{\circ}$ 

Table 1 Abiotic and biotic field parameters at the time of collection of Arenicola marina specimens at the sampling site in La Hume near Arcachon (Atlantic, France), mean values ± SD.

t1.2 t1.3	Sampling month	Salinity (‰)	T (°C) air	T (°C) tidal puddles	T (°C) in 20 cm depth	Length of tail shaft (cm)	Abundance $(m_{\perp}^{-2})$	Weight (g)
t1.4	August	$35.8 \pm 1.2$	$20.2 \pm 2.4$	$20.7 \pm 2.4$	$20.3 \pm 0.6$	$8.0\pm2.4$	$22.7 \pm 5.0$	$4.1 \pm 1.4$
	2005	n = 17	n = 13	n = 13	n = 13	n=6	n = 6	n = 29
t1.5	January	$26.7 \pm 0.5$	$7.6 \pm 2.4$	$9.5 \pm 2.3$	$7.1 \pm 0.8$	$13.0 \pm 2.4$	$23.0 \pm 9.1$	$4.7 \pm 1.6$
	2006	n = 25	n = 23	n = 23	n = 23	n = 10	n = 20	n = 72
t1.6	August	$34.1 \pm 0.6$	$22.1 \pm 2.4$	$25.8 \pm 4.1$	$22.2 \pm 0.8$	$12.9 \pm 2.9$	$28.4 \pm 11.2$	$5.7 \pm 1.8$
	2006	n = 20	n = 20	n = 20	n = 20	n=20	n = 20	n = 30
t1.7	February	$25.1 \pm 4.2$	$12.5 \pm 0.4$	$12.1 \pm 0.8$	$9.0 \pm 0.3$	$12.8 \pm 2.0$	$8.2 \pm 3.2$	$4.1 \pm 1.3$
	2007	n = 10	n = 10	n=10	n = 10	n = 10	n = 13	n = 30

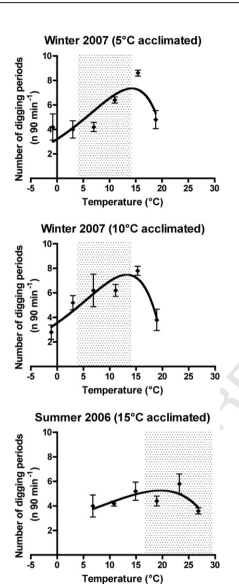


Fig. 1. Temperature dependent burrowing capacity in Arenicola marina. Mean values  $\pm$ SE for animals collected in summer 2006 and in winter 2007 at the French Atlantic coast, n=5. Winter animals were acclimated to 5 and 10 °C and tested between - 1.1 and 18.9 °C. Summer animals were acclimated to 15 °C and investigated between 6.8 and 26.8 °C (depicted from Schröer et al., 2009). Data were fitted to the equation  $EP(T) = F_1(T) + F_2(T) = (A_1e^{B1T} + C_1) + (A_2e^{B2T} + C_2)$  with EP(T) = temperature dependent muscle exercise performance capacity. The first term,  $F_1(T) = A_1e^{B1T} + C_1$ , represents the temperature dependence of aerobic processes supporting exercise performance. The second term,  $F_2(T) = A_2e^{B2T} + C_2$ , represents the parallel exponential rise in processes limiting aerobic scope and thus exercise performance capacity. For the winter data (5 °C acclimated):  $A_1 = -12.02$ ,  $B_1 = 0.1385$ ,  $C_1 = 1.800$ ,  $A_2 = 15.23$ ,  $B_2 = 0.1276$ ,  $C_2 = -1.807$ , r = 0.6472. For the winter data (10 °C acclimated):  $A_1 = 7.403, B_1 = 0.1173, C_1 = 0.9182, A_2 = -3.791, B_2 = 0.1494, C_2 = -0.9866, r = 0.6810.$ For the summer data:  $A_1 = 4.083$ ,  $B_1 = 0.06221$ ,  $C_1 = -0.4552$ ,  $A_2 = -1.081$ ,  $B_2 = 0.1038$ ,  $C_2 = 0.1652$ , r = 0.3723. Shaded area: naturally experienced habitat temperatures in the respective season.

temperatures. Temperature dependence was less pronounced and 332 lower maximum values were reached in summer animals, all values 333 stayed between 3.6 and 5.8 digging periods 90 min<sub>1</sub><sup>-1</sup>.

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#### 3.3. Respiration and ventilation experiment

Fig. 2 depicts temperature dependent weight specific volume flow 336 (ml  $g_{\perp}^{-1} h_{\perp}^{-1}$ ) in winter animals acclimated to 5 and 10 °C and in summer worms acclimated to 15 °C from two years. Specimens collected 338 in winter and acclimated to 5 °C showed an exponential increase in 339 weight specific volume flow over the experimental temperature 340 range with a  $Q_{10}$  value of  $5.57 \pm 0.63$  and remarkably larger error 341 bars at temperatures  $\geq 17.2$  °C.

Winter animals acclimated to 10 °C also exhibited an exponential 343 increase over the experimental temperature range, but the Q<sub>10</sub> of 344  $2.38 \pm 0.38$  was significantly lower than in 5 °C acclimated specimens. 345

Animals collected in summer 2005 and acclimated to 15 °C dis- 346 played a  $Q_{10}$  of 2.36  $\pm$  0.12 within the temperature range of exponential increase, similar to 10 °C acclimated winter specimens. At 2.8 °C 348 only one value was available, as the other worms stopped ventilation 349 activity during cooling. No statistics could be applied here. The value 350 recorded at 2.8 °C was lower than all values at 5.4 °C, therefore, the 351 one at 2.8 °C was excluded from the exponential range.

Summer lugworms from 2006 (also acclimated to 15 °C) displayed 353 the same exponential range as those from the preceding year, the 354 value at 2.9 °C differed significantly from the hypothetical value pre- 355 dicted by the regression curve. The  $Q_{10}$  of 2.24  $\pm$  0.23 was comparable 356 to the one evaluated in 2005.

In general, weight specific volume flow curves were similar in 358 winter and summer animals acclimated to 10 °C and 15 °C, respec- 359 tively. Only the data collected in winter animals acclimated to 5 °C 360 differed remarkably with around two times higher values in the tem- 361 perature range above 15 °C (Fig. 2).

Fig. 3 shows temperature dependent pumping frequencies (min<sup>-1</sup>) 363 during the phases of active ventilation. Winter specimens that were 364 acclimated at 5 °C showed a significant increase (F-test) over the 365 experimental temperature range. Values at 19.7 and 22.6 °C displayed 366 large variability with means significantly higher than in all other groups. 367 Animals acclimated to 10 °C in winter and to 15 °C in summer both 368 displayed a significant increase (F-test) with warming. Pumping 369 frequency in lugworms collected in summer 2005 did differed sig- 370 nificantly neither from the frequency recorded in winter animals 371 acclimated to 10 °C nor from data collected in summer 2006. The 372 only significant differences observed were the values at 10.9 and 373 23.2 °C from summer 2006 animals in comparison to both winter 374 groups.

Fig. 4 depicts the temperature dependent extraction coefficient 376 (%), i.e. the proportion of oxygen that the animals were able to withdraw from the inflowing water. Lugworms sampled in winter and 378 acclimated to 5 °C showed a more or less stable extraction coeffi- 379 cient in the temperature range between 0 and 5.1 °C while a slight 380 decrease was observed upon warming. In contrast, animals accli- 381 mated at 10 °C displayed a significant rise in the extraction coeffi- 382 cient from 0 to 17.7 °C followed by a slight decline. Summer specimens 383

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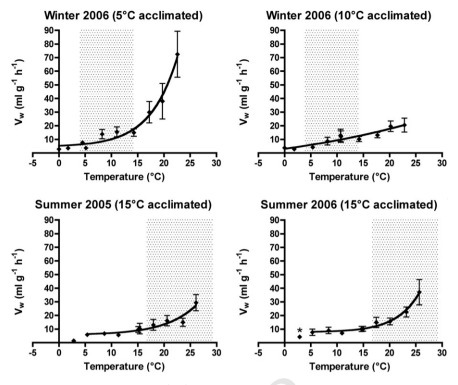


Fig. 2. Temperature (°C) dependent weight specific volume flow ( $ml g_{\perp}^{-1} h_{\perp}^{-1}$ , mean values  $\pm$  SE) in *Arenicola marina* dwelling in an artificial burrow. Animals were collected in summer 2005, in winter 2006 and in summer 2006 at the French Atlantic coast. n=5 except for winter (5 °C acclimated) at 22.6 °C, winter (10 °C acclimated) at 10.7 °C as well as summer 2005 at 15.4 and 5.4 °C with n=4 and summer 2005 at 2.8 °C with n=1. Winter worms were acclimated to 5 and 10 °C one half each, summer animals to 15 °C. The winter specimens were investigated from -0.0 to 22.8 °C, both summer groups were exposed to a temperature range from 2.8 to 26.1 °C. (Summer 2006 data depicted from Schröer et al., 2009). Data were fitted to  $V_w(T) = Ae^{BT} + C$  (Wittmann et al., 2008). For the winter 5 °C acclimated group: A=0.8083, B=0.1952, C=4.497, r=0.8118. For the winter 10 °C acclimated group: A=0.8083, B=0.1952, C=4.497, r=0.6989. For the summer 2006 group: A=0.08428, B=0.2273, C=7.832, r=0.7495. Asterisks (\*) designate data points, which are significantly different from the exponential regression curve (p=0.0234 for summer 2006 at 2.9 °C). Shaded area: naturally experienced habitat temperatures in the respective season.

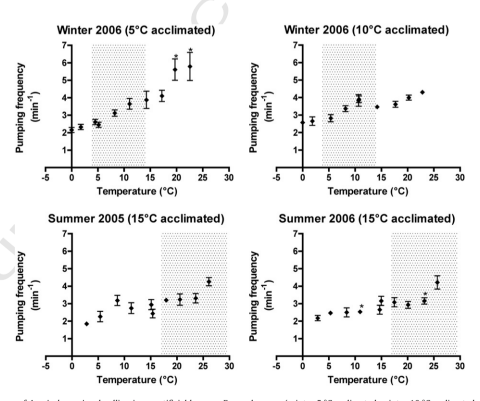
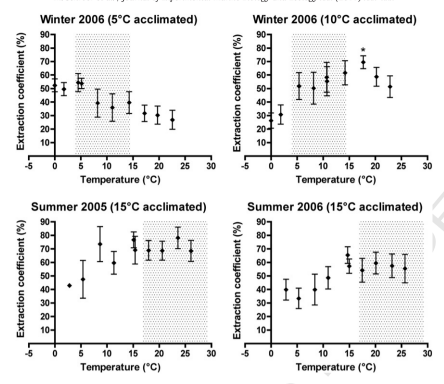


Fig. 3. Ventilatory performance of Arenicola marina dwelling in an artificial burrow. For each group (winter 5 °C acclimated, winter 10 °C acclimated, summer 2005 and summer 2006 (after Schröer et al., 2009)) pumping frequency (min $^{-1}$ ) is plotted against incubation temperature (°C). Mean values  $\pm$  SE; n = 5 except for winter (5 °C acclimated) at 19.7 and 22.6 °C, winter (10 °C acclimated) at 10.7 °C as well as summer 2005 at 15.4 and 5.4 °C with n = 4 and summer 2005 at 2.8 °C with n = 1. Sampling time and acclimation see legend Fig. 2. Incubation temperatures ranged from 2.8 to 26.1 °C for summer worms and from -0.0 to 22.8 °C for winter animals. Shaded area: naturally experienced habitat temperatures in the respective season.

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**Fig. 4.** Oxygen extraction by *Arenicola marina* dwelling in an artificial burrow. The oxygen extraction coefficient (%) versus incubation temperature (°C) is shown for the four groups (winter 5 °C acclimated, winter 10 °C acclimated, summer 2005, summer 2006 (from Schröer et al., 2009), mean values  $\pm$  SE, for further information see Fig. 2. \* = significantly higher than at -0.0 °C.

from both years showed a slight increase between 2.8 and 14.7 °C. With further warming, values stayed more or less constant. Again, the extraction coefficient in warm acclimated winter lugworms and summer specimens shows similar trends and differed from data obtained in winter animals acclimated at 5 °C.

## 3.4. Protein biosynthesis

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Fig. 5 shows the temperature dependent incorporation of <sup>13</sup>Cphenylalanine into proteins of the body wall after short and long term incubation times. A comparison of the three graphs reveals an overall increase of the amount of incorporated <sup>13</sup>C-phenylalanine with acclimation temperature. In particular, winter animals acclimated to 5 °C displayed no increase in <sup>13</sup>C-phenylalanine content over time at -0.9 °C. Data points for short and long incubation periods resulted similarly at around 4.4 nmol mg<sup>-1</sup> protein. At all other incubation temperatures, at least a slight increase between short and long incubation times was detectable with a significant difference at the highest incubation temperature (19.2 °C). In winter worms acclimated to 10 °C, the rise in the amount of incorporated amino acids over time was seen at all incubation temperatures. The maximum of  $24.26 \pm 1.55$  nmol mg<sup>-1</sup> protein was reached at 15.2 °C after long incubation times. Summer animals also showed an increasing incorporation of labelled phenylalanine over experimental time at all incubation temperatures. The largest amount of <sup>13</sup>C-phenylalanine was incorporated at 10.2 °C, reaching 32.48  $\pm$  1.24 nmol mg<sup>-1</sup> protein, a value significantly higher than at all other temperatures after short as well as long incubation periods.

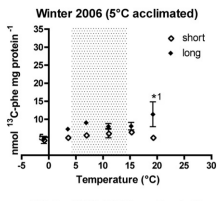
#### 4. Discussion

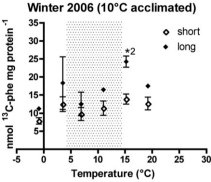
The aim of this study was to investigate potential seasonal acclimatisation effects in physiological performances of the lugworm *A. marina* from a southern population and to distinguish possible

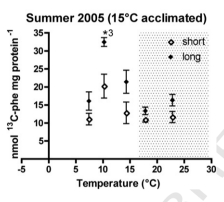
effects of acclimation temperature and inter-annual variability. Recently, we could show that animals from the southern-most population displayed a lower performance level than their counterparts from higher latitudes (Schröer et al., 2009).

Our results show clear seasonal differences in temperature tol- 418 erance and performance of A. marina. These are most likely due to 419 seasonal temperature changes, however, other biotic and abiotic 420 parameters also differ seasonally with potential influences on organ- 421 ismic performance. We observed a higher animal weight in summer 422 2006 compared to both winter samplings (Table 1), which might be 423 explained by a higher food availability in summer and more impor- 424 tantly, the onset of gamete production. In summer 2005, juveniles 425 were included in the samples and resulted in a lower average weight 426 than in 2006 and not higher than in winter. Under field conditions 427 growth in A. marina was only observed during spring and summer 428 until October (Beukema and de Vlas, 1979; Newell, 1948; Smidt, 1951; 429 Wolff and de Wolf, 1979) whereas during autumn and winter growth 430 was absent. Moreover, the weight of adults tends to decrease in winter 431 (Beukema and de Vlas, 1979; Newell, 1948). Pollack (1979) argued 432 that the lower mean animal weight in winter is not necessarily due 433 to individual weight loss but may also result from the immigration 434 of smaller worms to the sampling area and/or from a higher mortal- 435 ity among older and thereby larger worms. Any effects of lower 436 food availability in winter are exacerbated by a higher precipitation 437 in this season, easily visible in the changing salinity of the surface 438 water (Table 1). The lugworm shelters from the influence of fresh- 439 water by closing its burrow and reducing its pumping and feeding 440 activities during rainfall (Pollack, 1979). The maximally tolerated 441 daily salinity variation amounts to 4-6% (Amoureux, 1966).

Maximum food availability from April to October coincides perfectly 443 with the period of reproductive growth from the end of April until the 444 beginning of October (Cazaux, 1966). Although water temperature 445 has been well correlated with reproductive cycles and has often been 446 considered to be a major factor in their control, studies have also 447







**Fig. 5.** Amount of incorporated  $^{13}\text{C}$ -phenylalanine (nmol mg $^{-1}$  protein, means  $\pm$  SE) into protein of the cuticulo-muscular tube of lugworms, dependent on incubation temperature. Specimens were collected in August 2005 and January/February 2006. Winter worms were acclimated to 5 and 10 °C one half each and summer animals to 15 °C. Incubation temperatures ranged from -0.9 to 19.2 °C for the winter groups and from 7.4 to 22.8 °C for summer animals. Open diamonds: amount of incorporated  $^{13}\text{C}$ -phenylalanine after 30 to 120 min of incubation; closed diamonds: amount after 180 to 360 min; n = 4 for summer data except for short incubation times at 22.8 °C with n = 3, n = 3 for winter data. \*1 = significantly higher than the values after short incubation times at -0.9, 3.5 and 19.2 °C as well as at -0.9 °C after long incubation times. \*2 = significantly higher than the values after short incubation times at -0.9 and 6.9 °C. \*3 = significantly higher than all other values.

emphasised the importance of local food conditions (MacDonald and Thompson, 1986), as demonstrated by Newell et al. (1982) in *Mytilus edulis* populations, which experienced nearly identical temperature cycles but different regimes of food availability.

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460 461 Our measurements of acute somatic growth rates by tracking the incorporation of phenylalanine into the body wall revealed an optimum temperature for growth without substrate limitation at 15.2 °C in winter animals and at 10.2 °C in summer animals (Fig. 5). Interestingly, neither 15.2 °C is experienced in winter nor 10.2 °C in summer in the natural habitat. A look at the sea surface temperature (Ifremer, 2007), which correlates well with the sediment temperature in 20 cm depth (Nießing, 2006), showed that temperatures between 10 and 16 °C are experienced in March, April and May as well as in October, November and December. Our winter animals which were acclimated

to 10 °C showed maximum somatic growth at 15.2 °C and thereby, 462 the capacity to tolerate higher temperatures. This might indicate an 463 early shift to sustain spring conditions. Somatic growth is expected to take place in spring (March to May) when temperature is at its optimum and food is already available, as was the case in our aquarium 466 system. Phytoplanctonic and therewith phytobenthic growth takes 467 place a bit earlier in the year at Arcachon (Ifremer, 2007) compared 468 to the more northern beaches, for which the food availability data 469 are mentioned in the Introduction. The fact that <u>A</u>. marina juveniles 470 are found already at the end of March (Cazaux, 1966) also argues 471 for food availability already in March.

Our summer animals which were acclimated to 15 °C showed 473 maximum acute somatic growth at an exposure temperature of 10.2 °C. 474 Following the same rationale as before, this might be seen as a prepara- 475 tion for autumn conditions. This might also involve constraints on energy 476 budget in the warmth where cold exposure slows motor activity and 477 supports a shift of resources to growth. After spawning in autumn, the 478 worms are expected to replenish their glycogen reserves for winter 479 survival (see below). So growth is controlled by exposure rather 480 than acclimation temperature. Similarly complex interactions were 481 previously shown for juveniles of the oyster Ostrea edulis, which dis-482 played a maximum scope for growth at an acclimation temperature 483 of approximately 17 °C and an exposure temperature of approxi- 484 mately 25 °C, a common condition found in shallow waters during 485 the summer months (Buxton et al., 1981). In A. marina, all energy 486 is invested into reproductive growth during summer, while somatic 487 growth takes place in spring and autumn, suggesting a close rela- 488 tionship between energy available for growth, the reproductive 489 cycle and thermal limits. In fact, Füßner (2009) observed a higher 490 sensitivity to temperature changes in North Sea summer worms 491 compared to North Sea spring animals, which might be due to the 492 increased energy expenditure for reproductive growth.

Bayne and Newell (1983) suggested that scope for growth and 494 growth efficiency are more dependent on food availability than on 495 temperature. Both, a trade-off between growth and reproduction 496 and a strong dependence on food availability, were found in the 497 giant scallop Placopecten magellanicus: scope for growth was low 498 or negative during winter, rapid gamete maturation was observed 499 during the spring bloom and somatic weight declined during gamete 500 development while it increased after spawning and during periods 501 of low gametogenic activity (MacDonald and Thompson, 1986). 502 Similarly, low or negative growth during the winter and generally 503 higher values in the spring and/or summer are also described for 504 Chlamys islandica (Vahl, 1980), M. edulis (Bayne and Widdows, 1978; 505 Thompson, 1984) and Mya arenaria (Gilfillan et al., 1976). Whether 506 somatic tissue weight declines during gamete development in A. marina 507 has not been investigated so far and may depend on local temperature 508 and food conditions. For example, Chlamys varia shows both, simulta- 509 neous reproductive and somatic growth as well as gamete development 510 fuelled by somatic reserves, depending on environmental conditions 511 (Shafee, 1980). In Macoma balthica, somatic growth becomes negative 512 during gametogenesis when temperatures are high, but continues 513 despite gamete development at low temperatures (DeWilde, 1975). 514 High food availability combined with low temperatures and thereby a 515 reduced metabolic demand results in high scope for growth. These con- 516 ditions would allow for simultaneous somatic and reproductive growth 517 and might be found at Arcachon in April and May. In these months, 518 gamete development already begins (Cazaux, 1966) and temperature 519 ranges around the optimum for somatic growth (see above).

Laboratory experiments on somatic growth in *A. marina* were 521 already carried out by de Wilde and Berghuis (1979). Their study 522 was performed on animals from the Netherlands with constant 523 food supply and excluding the influence of reproduction by using 524 juvenile worms. In that study, maximum growth rates in length 525 and weight were observed at 20 °C with a small influence between 526 5 and 20 °C and a strong influence between 20 and 25 °C, as at 25 °C 527

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the growth rate was considerably lower than at 5 °C (de Wilde and Berghuis, 1979). It should be noted that these were acclimated, not acute growth rates. In natural populations of juvenile lugworms growth rates were usually found much lower than in the experiments. It may be assumed that the main limiting factor for growth in Arenicola is food limitation, caused either by food competition with other small mud flat inhabiting organisms or by a poor quantity and/or quality of the organic matter available (Boon and Haverkamp, 1979; de Wilde and Berghuis, 1979). Interestingly, the increase in biomass, i.e. the product of numerical density and mean weight, was highest at 10 °C in laboratory experiments, indicating a higher mortality with rising temperature, probably caused by permanently low oxygen concentrations induced by the combination of high amounts of organic matter and high temperatures (de Wilde and Berghuis, 1979). In field studies, the highest mortality rates were found from January to March, decreasing during spring and resulting in considerably lower values in summer than in winter (Pollack, 1979). This observation suggests once more that food availability is a limiting factor for growth in the field.

The only remarkable seasonal difference was found in exercise performance (Fig. 1). Corresponding to the concept of oxygen and capacity limited thermal tolerance (Pörtner and Knust, 2007; Pörtner et al., 2004), digging activity displays an asymmetric bell shaped curve, as already shown in a previous study (Schröer et al., 2009). This study demonstrated clear seasonal acclimatisation with a more pronounced optimum at lower temperatures in winter and a wider and less distinct performance curve with an optimum at higher temperatures in summer. The data suggest a trend for curve width to increase from winter to summer at the expense of a decrease in performance amplitude. Comparing cold- and warm-adapted lugworms from different latitudes (Schröer et al., 2009) a trade-off between the width and the amplitude of the performance curve also became apparent (cf. Angilletta et al., 2002; Huey and Hertz, 1984; Pörtner, 2006). For ectothermic vertebrates, it has already been shown that muscle twitch tension decreases with increasing temperature in fast-twitch muscles (Bennett, 1984). In addition, viscoelastic properties of cell membranes change with temperature, as seen for example in human red blood cells (Hochmuth et al., 1980). So higher temperatures in summer might result in a higher elasticity of cell membranes and change the worm's whole bodywall. For this reason, only a lower internal resting pressure (turgor) might be achieved in lugworms, causing lower amplitudes of body wall contraction pressure resulting in reduced digging performance. Seymour (1971) observed that a higher resting pressure resulted in a higher peak pressure in burrowing lugworms. Our observations of the internal pressure in burrowing lugworms by use of a catheter (data not shown) in fact exhibited a tendency of higher pressures at lower temperatures, but rising contraction frequencies with rising temperature. Therefore, the internal pressure might be the limiting factor at higher temperatures, while contraction frequency may be the limiting factor at lower temperatures.

Temperature fluctuation is higher in summer (16 to 29 °C) than in winter (4 to 14 °C), which might be one reason for a broader performance curve in summer. The lower performance amplitude in summer argues for a trade-off between reproductive growth and exercise performance. Performance curves of 5 and 10 °C acclimated specimens in winter were nearly identical leading to the conclusion that acclimation temperature alone has no effect on performance capacity. Consequently, seasonal acclimatisation does not only depend on temperature, but other seasonal changes (e.g. photoperiod, precipitation and food availability), as well as competing physiological processes like reproductive growth can also play an important role for the observed seasonal differences in muscular performance (i.e. digging activity).

Thermal acclimatisation of muscle performance has already been shown before in other marine invertebrates as in the European queen scallop, *Aequipecten opercularis* (Bailey and Johnston, 2005) and the giant scallop, P. magellanicus (Guderley et al., 2008). In these studies, 594 higher performance amplitudes also result in cold acclimated animals. 595 Cold acclimated A. opercularis attained higher swimming velocities 596 and accelerated faster at winter temperatures than warm acclimated 597 animals at summer temperatures (Bailey and Johnston, 2005). A 598 study by Guderley et al. (2008) reported that even handling stress 599 had less impact on cold than on warm acclimatised animals of the 600 species P. magellanicus. Guderley (2004) suggested that locomotor 601 performance and reproduction are closely coupled, as muscle meta- 602 bolic capacities fall in parallel with glycogen mobilisation for game- 603 togenesis, and reproductive fitness will be favoured more than 604 maintenance of performance. This interpretation might explain our 605 observations on lugworms as well. In our study, winter acclimatised 606 worms displayed the highest performance optima, as reproductive 607 growth takes place in summer (from end of April to the beginning 608 of October, Cazaux, 1966) and spawning occurs during autumn 609 (October to November). So in summer animals the gonads take up 610 most of the body mass, whereas winter animals do not invest any 611 energy into reproduction. Also A. opercularis and P. magellanicus 612 both have their spawning period until October and achieve perfor- 613 mance maxima in winter after spawning. Füßner (2009) observed a 614 much narrower thermal tolerance window in A. marina from the 615 North Sea during the time of spawning than in specimens during 616 the early stage of gamete production.

Consistent with previous findings (Schröer et al., 2009), perfor- 618 mance optima were found within the naturally experienced temper- 619 ature range during the respective season in this study. In winter, 620 modelled performance maxima of 14.2 and 13.3 °C, respectively, 621 were found close to the upper limit of naturally experienced habitat 622 temperatures of around 14 °C. In contrast, the modelled performance 623 maximum of 19.6 °C in summer was found close to the lower limit of 624 habitat temperatures during the respective season (around 17 °C). 625 This leads to the conclusion that performance, despite its seasonal 626 acclimatisation, is well adapted to the yearly mean habitat tempera- 627 ture of approximately 16 °C ensuring constant performance during 628 the whole year. By extrapolating the model curve, it also becomes 629 obvious that the upper critical temperature in summer, which is pre- 630 dicted by the model at 31.7 °C, nearly falls into the naturally experi- 631 enced range of habitat temperatures of up to 29 °C in summer. This 632 suggests that the population is more sensitive to warming effects in 633 summer than in winter.

Effects of temperature acclimation without the influence of reproduction have been investigated in tadpoles of *Limnodynastes peronii* 636 as well as tadpoles and adults of *Xenopus laevis* (Wilson and Franklin, 637 1999; Wilson et al., 2000). Cold acclimated organisms reached a higher 638 swimming velocity at low temperatures than warm acclimated specifies, while at high temperatures warm acclimated animals showed a 640 higher swimming performance than cold acclimated ones. This simple 641 relationship is overlaid with the effects of reproductive growth in our 642 study when comparing digging activity in winter and summer acclimated lugworms.

Despite a nearly identical performance curve, the group of winter 645 animals acclimated to 5 °C showed a much higher volume flow than 646 those acclimated to 10 °C. Differences became obvious especially at 647 temperatures above 15 °C, resulting in a high  $Q_{10}$  value of  $5.57 \pm 0.63$  648 compared to values between 2.24 and 2.38 in the other winter and 649 summer groups. 5 °C acclimated winter specimens showed their 650 highest extraction coefficients at temperatures from 0.0 to 5.1 °C, 651 whereas 10 °C acclimated winter worms and both groups of sum-652 mer worms exhibited their lowest extraction coefficient values in 653 this temperature range. Cold compensation at the cellular level like-654 ly occurred at 5 °C. Sommer and Pörtner (2004) found changes in 655 mitochondrial functions taking place in A. marina from the North 656 Sea, which were acclimated to 0 °C in comparison to those acclimated ed to 5 and 11 °C. These findings suggest that cold acclimatisation 658 occurs below a threshold temperature rather than progressively 659

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with falling environmental temperatures. The other way round, returning to warm acclimatisation also seems to occur stepwise, as Wittmann (2005) found different thermal tolerance windows in worms which experienced a warming pulse of 3 °C compared to those which were investigated before. Somatic growth (Fig. 5) was minimal in our 5 °C acclimated compared to 10 °C acclimated winter lugworms. These findings could be an indication for a temperature induced dormant condition as it is found in the common shallowwater and littoral bivalve Cardium (= Cerastoderma) edule L. in winter (Newell and Bayne, 1980). Dormancy so been found in other marine polychaetes like *Lanice conchilego* eres, 1997). Dormant individual relies on carbohydrate reserves for maintenance energy requirements (Newell and Bayne, 1980). Lugworms also show a decrease in the glycogen content of the body wall between November and February (Nießing, 2006), which coincides with the time of low food availability (see above). Accordingly, winter specimens in our study displayed a significantly lower fresh weight than summer animals (Table 1). Field observations revealed a lower faeces production and hence food uptake from October to January compared to the spring and summer months (Pollack, 1979), possibly initiated by a drop in temperature and/or a reduced food availability. It was shown in laboratory experiments that faeces production and hence feeding activity is dependent on the food content of the sediment. Considerable differences between actual numbers of lugworms and number of faecal casts occurred, indicating part of the animals to be inactive at less favourable conditions in poor sediments (de Wilde and Berghuis, 1979).

Also the organic matter content of the water was shown to influence the pumping activity (Krüger, 1964). Our 5 °C cold-acclimated winter lugworms pursued the strategy of a "conserver", which lives under conditions of short food, feeds at a low rate because a higher rate might increase the costs of foraging more than the gains, and has a low growth rate in favour of longevity (Branch et al., 1988). In contrast to volume flow, pumping frequency, extraction coefficient and growth, the modelled exercise performance curves and maxima (Fig. 1) were nearly identical for both winter groups. This emphasises that maintenance of muscle exercise capacity seemed to be important for winter lugworms. An explanation might be the observation of active winter migrations in A. marina, initiated by temperatures close to critical values (Werner, 1956). When the limits of acclimatory adjustment are reached, the animals leave their habitat and rebury in a lower zone of the intertidal. De Wilde and Berghuis (1979) made a similar observation in a laboratory experiment, when lugworms fed and maintained at 5 °C left the sediment for migration.

## 5. Conclusions

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Seasonal differences become obvious in exercise performance curves of A. marina. Winter animals exhibit an optimum at around 14 °C and a high performance amplitude. In summer, the optimum is shifted towards 19 °C, accompanied by a widening of the performance window and lower performance amplitudes. A trade-off between exercise capacity and reproductive growth seems to take place in summer. In addition to temperature food availability is likely an important factor controlling seasonal acclimatisation processes. Somatic growth may occur mostly in spring and autumn, when food is available, and outside the reproductive period. The experimentally determined optimum combination of acclimation and exposure temperatures for growth matches well with the temperature conditions found in spring and autumn in the worms' natural habitat. Altogether, the present results show that temperature determines the metabolic state of lugworms despite of the season. Furthermore, our data confirm that the investigated lugworm population from Arcachon lives at the upper level of its thermal limit, making it most susceptible for warming waters in the near future.

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