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3	Geographical variation in thermal tolerance within Southern Ocean marine ectotherms.
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13	Short title: geographic thermal tolerance
14	
15	Abstract
16	Latitudinal comparisons of the Southern Ocean limpet, Nacella concinna, and clam, Laternula
17	elliptica, acclimated to 0.0°C, were used to assess differences in thermal response to two
18	regimes, 0.0, 5.1 to 10.0°C and 2.5, 7.5 to 12.5°C, raised at 5.0°C per week. At each temperature,
19	tissue energy status was measured through a combination of O ₂ consumption, intracellular pH,
20	cCO2, citrate synthase (CS) activity, organic acids (succinate, acetate, propionate), andelylates
21	(ATP, ADP, AMP, ITP, PLA (phospho-L-arginine)) and heart rate. L. elliptica from Signy
22	(60°S) and Rothera (67°S) had the same lethal limits (7.5-10.0°C), but small yet consistent
23	indicators (O2, CS activity, PLA, ATP, organic acids and cCO ₂) suggest that Rothera L. elliptica
24	had lower critical and pejus (=getting worse) limits than Signy L. elliptica. N. concinna, which
25	experiences a wider thermal regime, had higher lethal limits (10.0-12.5°C). However, at their
26	Northern geographic limit N. concinna living in a warmer environment (South Georgia, 54°S),
27	had a lower critical limit (5.1-10.0°C; O2, PLA and organic acids) than Rothera and Signy N.
28	concinna (10.0-12.5°C). This lower limit indicates that South Georgia N. concinna have different
29	biochemical responses to temperatures close to their thermal limit, which may make them more
30	vulnerable to future warming trends.
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Key words: *Nacella concinna*; *Leternula elliptica*; Southern Ocean; tissue biochemistry; *pejus*limits; critical limits; lethal limits; latitudinal comparisons;

34

35 **1. Introduction**

36 Applying the principles of symmorphosis (Weibel, 2000) to the oxygen supply cascade 37 provides a conceptual framework to explain how temperature affects oxygen availability to 38 tissues, and hence tissue energy status. This has developed towards a unifying principle that the 39 physiological thermal limits of aquatic ectotherms are defined by oxygen availability, the 40 balance between supply and demand (Pörtner, 2002; Pörtner, 2006; Peck et al., 2007). Within 41 their optimal range ectotherms are expected to have the physiological flexibility to cope with 42 normally experienced seasonal variation and tissue energy status should be maximal. However, 43 in the face of extreme conditions, or towards range limits, aerobic scope is reduced, as more of 44 the available energy is required to cope with routine metabolic processes and oxygen supply 45 mechanisms become sub-optimal. There is growing evidence that the effects of thermal stress act 46 hierarchically, from high to low complexity, i.e. whole body processes are affected before 47 biochemical reactions and cells (Pörtner, 2002; Pörtner and Knust, 2007; Pörtner et al., 2007). 48 Improving our understanding of how functionality is lost and energy is partitioned between 49 processes will improve our understanding of the likely impact of climate change on species 50 distributions and resulting ecosystem shifts.

51 Thermal sensitivity can be assessed at a number of levels from whole animal activity 52 through to tissue energy status and finally mortality. Characterising changes in tissue 53 biochemistry has been developed as a test for *critical* thermal limits in aquatic ectotherms which 54 indicate the transition to anaerobic metabolism (cf Pörtner, 2002). Within the window bounded 55 by critical temperatures, an even narrower window is bordered by pejus limits (pejus: getting 56 worse) which indicate when temperatures move beyond the optimal range and functionality starts 57 to be lost, which is paralleled by a shortfall in oxygen supply and demand until critical limits are 58 reached and anaerobic metabolic pathways are recruited. Species tested this way include the 59 boreal freshwater gadoid, Lota lota (Hardewig et al., 2004), the worms, Sipunculus nudus 60 (Zielinski and Pörtner, 1996), several populations of Arenicola marina (Sommer et al., 1997), 61 and several Antarctic species (e.g. Limopsis marionensis; Pörtner et al., 1999) including both L. 62 elliptica (Peck et al., 2002) and N. concinna (Pörtner et al., 1999). Biochemical parameters were

63 selected that measure the integrity of cellular systems and identify the temperature at which there is a departure from homeostasis (pHi and cCO₂), with the main focus on those that measure the 64 65 aerobic status of the animal, circulation (heart rate), aerobic metabolism (O₂ consumption and citrate synthase activity), tissue energy status (balance of adenylates) and recruitment of 66 67 anaerobic pathways (build up of organic acids). We aim to utilise these techniques to compare 68 pejus limits, critical and lethal limits of N. concinna and L. elliptica from different locations 69 within the Antarctic. These measures will be used to look for adaptive differences between 70 locations that relate to experienced thermal environment.

71 We exploit access to a unique latitudinal gradient within the Southern Ocean, Rothera 72 (67°S), Signy (60°S) and South Georgia, (54°S) which enable us to separate latitude and 73 temperature (Barnes et al. 2006). Our highest latitude site at Rothera Point, Adelaide Island, has 74 an annual seawater temperature ranging from -2° C in winter to $+1^{\circ}$ C in the height of summer (at 75 15m depth; Barnes et al., 2006). Signy Island, within the South Orkney Islands (60°C) is on the 76 edge of the Weddell Sea gyre and despite its more northerly location has the same annual 77 seawater temperature range to that of Rothera (Clarke and Leakey, 1996). However the sub-78 Antarctic Island of South Georgia (54°S), although still within the Southern Ocean (south of the 79 Antarctic circum-polar current) and therefore the biological Antarctic (Barnes et al., 2005), has 80 almost twice the annual range due to a higher summer maximum (-1 to $+4^{\circ}C$ at 10-20m; Barnes 81 et al., 2006). In response to the constant cold at Rothera benthic marine ectotherms are generally 82 stenothermal, they have a limited temperature range, upper lethal limits typically occurring 83 between 5 and 10°C (Peck, 2005) but for some species are considerably lower than these, e.g. the 84 brittle star *Ophionotus victoriae* which cannot survive even 1 month at 2°C (Peck et al, *in press*). 85 Furthermore, limits for critical activities are often considerably lower than the lethal limits (2 to 86 4°C; Peck et al., 2004). The +4°C maximum summer seawater temperature at South Georgia 87 therefore has the potential to impact these species, unless they have adaptive capacity to cope 88 with this thermal range. It is interesting to note that the range of O. victoriae does not stretch as 89 far as South Georgia.

Symmorphosis (Weibel, 2000) and the climate variability hypothesis (Stevens, 1989)
both predict that species, which experience a wider range of temperature, will have greater
thermal flexibility. From this, sub-littoral species living at South Georgia should be able to cope
with higher temperatures than the same species from Rothera. Habitat temperatures would also

94 predict that the thermal response of individuals from Signy would be closer to those from 95 Rothera than South Georgia. Although littoral species will experience a much higher range of 96 temperatures during low water emersion, habitat temperatures still predict the warmest 97 environment at South Georgia. The highest recorded littoral temperature is 12.3°C at Rothera 98 (Waller et al., 2006), 10.7°C at Signy (Barnes et al., 1996) and 15.8°C at South Georgia 99 (Davenport, 1997). The littoral limpet, Nacella concinna (Strebel 1908), would therefore be 100 expected to have a higher thermal capacity than the sub-littoral clam, Laternula elliptica (King 101 and Broderip 1831) and higher thermal limits in N. concinna and L. elliptica would be predicted 102 from South Georgia, than Signy and Rothera. However, species living at their range limit, where 103 they experience temperatures close to their maxima may have a reduced ability to cope with 104 further temperature increases (Tomanek, 2005; Deutsch et al., 2008); adaptations that enable 105 ectotherms to live in warmer environments may occur at the expense of further acclimatory 106 capacity (Stillman, 2003).

107 Through sampling these two species from 3 locations within the Southern Ocean, at the 108 same time of year, we aim to conduct the first latitudinal comparison of thermal tolerance within 109 the Southern Ocean. Environmental history determines thermal response (e.g., Osovitz and 110 Hofmann, 2005) and so all animals were acclimated to the same temperature (0.0°C) and 111 maintained under as near identical conditions as possible for 2 months before thermal responses 112 were tested. Any differences in thermal response should therefore indicate adaptive differences 113 between locations.

114

115 **2. Materials and Methods**

116 2.1. Collection and culture

Limpets, *Nacella concinna* and the soft shelled clam, *Laternula elliptica* were collected
during cruise JR109 on the RRS James Clark Ross and concurrently at Rothera, the British
Antarctic Survey Research Station at Adeleide Island on Antarctic Peninsula, during early
summer as detailed in table 1. Except for *N. concinna* from Signy, which were collected from the
lowest part of the intertidal, all animals were collected by SCUBA divers.
After collection all animals were maintained in recirculating aquaria at 0.0±0.3°C, for 63

123 days until the start of experiments. Water quality was maintained through biological filtration,

124 U.V. sterilisation, protein skimmers and daily water changes. During this period animals were

not fed to reduce any metabolic increment due to feeding (SDA), which can last up to 25 days in
Antarctic benthic ectotherms (Peck, 1998). Nonetheless, *N. concinna* were observed grazing on
biofilms on the tank walls and *L. elliptica* were seen with siphons open, so both species are likely
to have continued low level feeding.

129 At the start of incubation experiments animals were transferred to two 200 litre jacketed 130 acrylic tanks (Engineering Design and Plastics, Cambridge, Cambs.) attached to LTD20G 131 thermocirculators (Grant instruments Ltd, Shepreth, Cambs.), which maintained temperatures to 132 an accuracy of $\pm 0.2^{\circ}$ C. The method was similar to that that used by Morley et al. (2007) and 133 Peck et al. (2002; 2004; 2007 and 2008). Animals were allowed 48 hours to recover from 134 movement stress before temperatures were altered or measurements taken. Temperature was then 135 raised at approximately 0.1°C per hour until the required temperatures were reached and then 136 animals were allowed 48 hours to recover from any metabolic overshoot (Peck et al., 2002). Due 137 to time constraints animals were separated into two groups, which were taken through different 138 5.0°C temperature steps, tank 1) 0.0, 5.1 and 10.0°C, tank 2) 2.5, 7.5 and 12.5°C. Following this 139 protocol animals were tested over approximately 3 weeks in each tank.

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141 2.2. Oxygen consumption

142 Routine metabolic rate was measured in closed cell respirometers, following the 143 methodology of Peck (1989), except that oxygen concentration was measured with a Fibox-3 144 oxygen meter (Presens GmbH, Regensberg, Germany; e.g. Morley et al., 2007). Oxygen 145 sensitive foils were calibrated before each measurement using 5% w/w sodium diothionite for 146 0% and fully aerated water for 100%. During trials oxygen concentration was not allowed to fall 147 below 70% of air saturation, which is above the threshold for oxy-regulation for L. elliptica 148 (Peck et al., 2002). Two empty chambers (controls) were run with each trial to account for 149 background oxygen consumption, which was routinely less than 10% of the animal's 150 consumption. The basal metabolic rate of 4-5 individuals was measured for three consecutive 151 days after which data were analysed for significant differences in oxygen consumption between 152 days. When significant differences were found, daily measurements continued until three stable 153 readings were obtained and then temperature was raised. The volume of each animal was 154 measured through displacement and subtracted from the volume of water in the respirometer. At 155 the end of the experiment dry weight (constant weights at 60° C) and ash free dry weight (dry

weight minus the weight of ash remaining after ignition at 475°C for 24h) of each whole animal
was measured.

158 At each temperature the circulatory system of L. elliptica was monitored with the 159 impedance-based heart rate monitoring system (Buchan et al., 1988). Impedance systems record 160 the frequency of heart rate through variations in resistance between two electrodes. Methods 161 used were similar to those of Peck et al. (2002). Two 1-mm holes were drilled through the shell 162 either side of the heart, and 2mm of tinned 18 standard gauge copper wire electrodes, with the 163 coating removed from the tips, were inserted and glued in place with cyanocrylate gel adhesive. 164 4 Signy and 4 Rothera L. elliptica had electrodes implanted in each of the two temperature tanks. 165 At the end of each temperature step the foot muscle from 4 animals was freeze clamped 166 in liquid nitrogen and stored at -80°C for biochemical analysis at the Alfred Wegener Institute in 167 Germany (except South Georgia and Signy N. concinna at 10°C for which only 3 animals were 168 available). Foot muscle was the chosen tissue as it performs a similar function in both N. 169 concinna and L. elliptica. Tissue was ground in a pestle and mortar, under liquid nitrogen, and a 170 number of extracts produced for biochemical analysis.

Temperature steps continued until 50% mortality occurred in the tanks. A further set of
20 *N. concinna* and *L. elliptica* from each location was maintained in the flow through aquarium,
at ambient temperature, for the duration of the experiment, with no mortalities.

174

175 2.3. Biochemical analysis

176 Intracellular pH (pH_i) and cCO_2 could only be measured in tissues of L. elliptica as N. 177 concinna foot muscle has large calcium carbonate stores, which would interfere with 178 homogenate acidity (Pörtner et al., 1999). Extracts were prepared in a media containing 160mmol.l⁻¹ potassium fluoride and 1.0 mmol.l⁻¹ nitrilotriacetic acid following the methods of 179 180 Pörtner et al. (1990). pH_i was measured in the supernatant of the centrifuged homogenates at 181 animal incubation temperature, using needle type fibre optic pH sensors attached to a pH1 micro 182 pH meter (Presens GmbH. Regensburg, Germany) whose analogue output signal was boosted 183 and digitised through a Powerlab system (AD instruments). Due to the high sensitivity of the optode system to solution ionic strength, KCl was added to each calibration buffer to match L. 184 *elliptica* tissue ionic strength of 0.16mol L^{-1} . Due to the small change in tissue pH, calibration 185 186 was conducted using three buffers covering the pH range 6.865 to 7.413 (at 25°C) whose pH

187 change with temperature had already been calculated. pH sensors were re-calibrated using188 buffers cooled to each incubation temperature.

To measure cCO_2 100µl of this supernatant extract was injected into a 10ml sealed sample vial, containing 1.5ml of nitrogen gassed HCl (0.05M). The sample was rotated for 10 minutes and then 50µl of this gas phase was injected into an Agilent technologies 6890N Network gas chromatogram through a G1888 network headspace sampler. cCO_2 was measured with a HP-PLOTQ column and a TCD detector. Triplicate peak areas were converted to cCO_2 by plotting a calibration curve for standard samples between 0.5 mM and 10.0 mM dissolved inorganic carbon ($r^2 = 0.998$).

196 Tissue citrate synthase activity was measured from tissue extract at a ratio of 1g of tissue 197 to 9ml of extraction buffer, following the methods of Sidell et al. (1987). Enzyme activity was 198 measured at animal incubation temperature with a Beckman UV-DU 7400 spectrophometer 199 fitted with a temperature controlled cuvette holder and diode array detector.

200 To measure both the build up of the end products of anaerobic metabolism (organic 201 acids) and tissue energy status (the adenylates) tissue homogenate was prepared using 0.6 mol.l⁻¹ 202 perchloric acid (PCA) (modified from Hardewig et al., 1998). Diluted PCA extracts were 203 injected into an ICS 2000 ion chromatography system fitted with a Gilson Diluter 401 204 autosampler. Organic acids were separated on an ion exclusion column (Dionex ICE-AS 1) at 205 60°C fitted with an AMMS-ICE 4mm micro-membrane suppressor cell, with 0.20mM heptafluorobutyric acid (HFBA) as eluent at a flow rate of 1 ml.min⁻¹ and 5mM 206 207 tetrabutylammonium hydroxide (TBAOH) as the regenerant. The signal was detected on a DS6 208 conductivity detector. Duplicate sample peaks were converted into concentrations using calibration curves with standard solutions, succinate (12 μ M to 212 μ M, r² =0.997) acetate 209 $(14\mu M \text{ to } 296\mu M, r^2 = 0.997)$ and propionate $(12 \mu M \text{ to } 189\mu M, r^2 = 0.999)$. 210 211 Adenylates were measured on a Beckman PACE/MDQ capillary electropheris system 212 following the adapted methods of Casey et al. (1999). The capillary was uncoated fused silica, 213 50µm ID, 50 cm long. The pH of homogenates was set to 9.5, diluted in separation buffer 214 (40mM tetraborate buffer and 10mM sodium chloride (pH 9.7)) and 50µl of 4 mM uric acid was 215 added to 200µl of each standard and sample as an internal standard. The absorbance signals of 216 separation were simultaneously detected at different wavelengths with a photo diode array 217 detector: at 254 nm, adenosine, AMP, ATP, ADP and ITP; at 200 nm, arginine and PLA; at 290

218 nm, internal standard uric acid. Duplicate peaks were converted into concentrations using the

following calibration curves: adenosine (71.5 μ M to 572 μ M, r² =0.995), AMP (70.5 μ M to 564

220 μ M, r² =0.988), ATP (80.4 μ M to 643 μ M, r² =0.995), ADP (71.3 μ M to 570 μ M, r² =0.966),

221 ITP (70.2 μ M to 561 μ M, r² =0.992), arginine (82.0 μ M to 656 μ M, r² =0.986) and PLA (68.8

- 222 μ M to 550 μ M, r² =0.996).
- 223

224 2.4. Analysis

225 Conducting the experiment in two tanks simultaneously created a mixed design where, 226 within tanks, some individuals were sampled more than once (e.g. for oxygen consumption), but different individuals were sampled between tanks. As there was a small, non significant, effect of 227 228 individual (as a random factor in a GLM analysis) a simple analyses was conducted, ignoring repeated observations on some individuals (P. Rothery, pers. coms.). ANOVAS with Tukev post-229 230 hoc tests were used to test for differences between locations and temperatures for each species. 231 Where ANOVAS indicated only significant differences between locations a separate analysis 232 was conducted for each location. Significant differences was accepted as P < 0.05 throughout. 233 Oxygen consumption was standardised to that of a 0.373g dry mass N. concinna and a 10.6g dry 234 mass L. elliptica (the mean mass of animal used) using scaling coefficients of 0.82 (Fraser et al., 235 2002) and 0.73 (Peck et al., 2002) respectively.

236

237 **3. Results**

238 3.1. Animal size

The shell length of *L. elliptica* sampled for biochemistry was not significantly different between locations, table 2. However, due to logistic constraints on collections by SCUBA divers, limpets from Rothera were smaller than those from Signy and South Georgia. *N. concinna* collected inter-tidally from Signy were not significantly different in size than *N. concinna* collected sub-tidally from South Georgia.

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245	3.2. L	. elliptica
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More than 50% of *L. elliptica* died during the transition from 5.0 to 10°C and from 7.5 to 12.5°C so the experiment was stopped at this stage and the upper thermal limit recorded as between 7.5 and 10.0°C. Although the sample size was too small to detect significant effects, heart rate increased with temperature between 0.0 and 5.1°C. 50% of animals with implanted
impedance wires died at 7.5°C so heart rate was not measured at this temperature.

251 Temperature had a significant effect on every biochemical parameter, there were a few 252 significant differences between Rothera and Signy L. elliptica (Table 3) and some consistent 253 patterns that indicate differences between L. elliptica from the two locations. Although generally 254 not significant, at 0.0°C, Rothera L. elliptica had higher oxygen consumption, CS activity and 255 higher concentrations of high energy phosphates (PLA, ITP and ATP) than Signy L. elliptica 256 suggesting an overall higher metabolism at 0.0°C. Between 0.0 and 2.5°C there was also a, non-257 significant, but consistently greater magnitude change in PLA and ATP (reduction), cCO₂ 258 (increase) and organic acids (increase of succinate and acetate) in Rothera than Signy L. elliptica

(Fig 1, Table 3).

The oxygen consumption of *L. elliptica* rose between 0.0 and 2.5°C for specimens from both sites, but no oxygen consumption was significantly different from that at 0.0°C. There was, however, a significant drop in oxygen consumption between 2.5 and 7.5°C for both Signy and Rothera *L. elliptica*. Citrate synthase activity at 2.5, 5.1 and 7.5°C was not significantly different from that at 0.0°C. However, in a similar fashion to oxygen consumption, *L. elliptica* from both locations had a peak in enzyme activity at 2.5°C, which was significantly higher than activity at both 5.1 and 7.5°C.

267 The energy status of the high energy phosphates fell as the concentration of their low energy forms, arginine, AMP and ADP, increased at 7.5°C in L. elliptica from both locations. 268 269 PLA the buffer that allows rapid rephosphorylation of ADP yielding ATP was also reduced at 270 7.5°C. The response of the triphosphates, ITP and ATP, to temperature was slightly different 271 between L. elliptica from Signy and Rothera. The ATP levels of Rothera L. elliptica were lower 272 at both 5.1 and 7.5°C than at 0.0°C, whilst the ATP levels of Signy L. elliptica were only 273 reduced at 7.5°C. The ITP concentration of Rothera L. elliptica generally fell upon warming but 274 was only significantly lower than 0.0°C at 7.5°C, whilst the ITP concentration of Signy L. 275 elliptica was lower at both 0.0°C and 7.5°C than 5.1°C. 276 The concentration of all three organic acids, succinate, acetate and propionate was

elevated at 2.5°C in Rothera *L. elliptica* and at 5.0°C in both, but was only significantly higher at
7.5°C. There were no significant differences in acetate and propionate accumulation patterns

between locations except that succinate rose to a significantly higher level in Signy than Rothera

280 *L. elliptica* at 7.5°C.

Intracellular pH was significantly higher at 0°C than at 2.5, 5.1 or 7.5°C for *L. elliptica* from both locations. There was also a significant drop in pHi between 5.1 and 7.5°C. Tissue CO_2 was highly variable and there was only one significantly elevated value, in Rothera *L. elliptica* at 5.1°C.

285

286 *3.3. N. concinna*

287 More than 50% of *N. concinna* died during the transition from 7.5 to 12.5°C and 10.0 to 288 15.0° C so the experiment was stopped at this point. There was no significant effect of location or 289 temperature on arginine but there were complex, significant effects on the other adenylates, with 290 higher concentrations of both low (AMP and ADP) and high (ATP) energy adenylates at 10.0°C 291 (Fig. 2). At 10.0°C AMP was significantly higher in Rothera N. concinna than all treatments 292 except Signy N. concinna at 2.5°C and South Georgia at 10.0°C. At 10.0°C ADP was higher in 293 *N. concinna* from all locations. Although there was no significant difference in ATP 294 concentration from that at 0.0°C, at 10.0°C the concentration of ATP was higher at all locations 295 than at 2.5°C. However, despite this significant increase in ATP for all locations, the increase for 296 South Georgia N. concinna appeared to be less (Fig. 2). The PLA concentration of South Georgia 297 *N. concinna* was also lower at 10.0°C than at either 0.0 or 5.1°C. 298 Despite a general increase in the organic acids with increasing temperature, succinate,

acetate and proprionate were only significantly elevated in South Georgia *N. concinna* at 10.0° C (Fig.3). The large inter-individual variation in organic acids at 10° C is typical of biochemical responses around physiological limits as some individuals reach their limits before others. For *N. concinna* from all locations, citrate synthase activity determined in tissue samples collected at each temperature generally increased upon warming but was only significantly higher at 7.5 and 10.0° C (Fig.4).

- 305 Oxygen consumption was very different between *N. concinna* from different locations.
 306 Oxygen consumption increased in a linear fashion with temperature (T) for Rothera (O₂cons =
- 0.14T + 1.58, R²=0.27, p<0.05, F=6.8) and Signy (O₂cons = 0.10T + 0.92, R²=0.57, p<0.01, p<0.01)
- 308 F=22.9) *N. concinna* (Fig 4). Oxygen consumption for South Georgia *N. concinna* reached a
- 309 similar maximum value but was best described by a curvilinear relationship ($O_2 cons = 0.64T$

 $0.06T^2 + 0.76$, R²=0.45, p<0.01, F=6.6). The calculated maximum (the point of inflexion of the differential equation) occurred at 5.3°C, but the highest oxygen consumption was recorded at 2.5°C (Fig. 4).

313

314 4. Discussion

Measuring a suite of physiological and biochemical parameters gave a comprehensive picture of how tissue energy status changed in response to a 5.0°C per week temperature elevation. It also allowed comparisons of the responses of *L. elliptica*, and *N. concinna*, collected from different geographic locations. As all animals were collected at the same time, were held under nearly identical conditions for approximately 2 months before thermal sensitivity was tested and they were tested over a maximum of 3 weeks, any consistent differences between locations are most likely due differences in response between animals from these locations.

322

4.1. L. elliptica

324 The critical limit, which was confirmed by a drop in oxygen consumption, a drop in 325 energy status of the adenylates, an increase in anaerobic end products of metabolism (the organic 326 acids) and a drop in intracellular pH, occurred between 5.1 and 7.5°C, which is comparable to 327 that measured previously, over a similar rate of temperature change (6° C; Peck et al., 2002). 328 Anaerobic metabolism is generally considered one of the protective mechanisms that allow time-329 limited survival above critical temperatures (Pörtner, 2006). However, molluscs in particular can 330 have high anaerobic capacities that can be utilised to survive hypoxia for days or weeks, 331 particularly when accompanied by metabolic depression (e.g. Holmes and Miller, 2006; Morley 332 et al., 2007; Long et al., 2008). Reductions in citrate synthase activity and metabolic rate suggest 333 that thermal limitation may have started at a lower temperature, with *pejus* limits below 5.1°C. 334 Succinate and volatile fatty acid formation set in somewhat earlier in Rothera than the Signy L. 335 *elliptica*. This would indicate that, as a consequence of severe oxygen limitation, the critical 336 temperature is lower in the Rothera than the Signy population. The greater, but non-significant, 337 biochemical changes between 0.0 and 2.5 °C for Rothera L. elliptica, coupled with significant 338 accumulation of carbon dioxide in the haemolymph, which shows that Rothera L. elliptica have a 339 lower capacity for gas exchange at 5.0 °C (cf Peck et al., 2002), also suggest that pejus limits are 340 lower in Rothera than Signy L. elliptica. After 2 months at 0.0 °C, there were also signs of

differences in cold acclimation of; CS activity, oxygen consumption and the turnover of high
energy phosphates were consistently (but not significantly) higher in Rothera than Signy *L*. *elliptica* at 0.0 °C.

344 The lethal limit in the current study also occurred at a similar temperature, between 7.5 345 and 10.0°C (within 15 days), to that recorded previously (9°C within 25 days; Peck et al., 2002). 346 The similarity of lethal limits between Rothera and Signy L. elliptica, is consistent with the 347 hypothesis that thermal capacities will be matched to the experienced environment (both -2 to 348 +1°C), although the lower *pejus* limits of Rothera L. elliptica compared to Signy indicate subtle 349 differences in thermal tolerance between locations. Clearly, studies incorporating L. elliptica 350 from South Georgia, which is at the northern limit of its range, are needed to quantify how close 351 it is living to its upper thermal limits and therefore with reduced capacity to cope with further 352 temperature increases (Stillman, 2003). Such studies are crucial to our understanding of species 353 wide thermal capacity and therefore the ability of species to cope with environmental change. 354 Acclimations of *L. elliptica* at 3-4°C would indicate if South Georgia *L. elliptica* have 355 adaptations to cope with higher maximum experienced summer temperature.

356

357 *4.2. N. concinna*

358 Due to the constraints on geographical sampling within the Southern Ocean, there were 359 sampling differences between N. concinna collected at different locations. N. concinna were 360 collected from the bottom of the inter-tidal from Signy, compared to sub-tidal animals collected 361 from South Georgia and Rothera, however, all N. concinna were collected from depths shallower 362 than 6m. A recent study found small, non-consistent, genetic differences between inter-tidal and 363 15m N. concinna from Potter Cove, King George Island, but not between inter-tidal and 6m N. 364 concinna (de Aranzamendi et al., 2008). Different biochemical responses to emersion were also 365 evident between 15m and intertidal N. concinna, also from King George Island (Weihe and 366 Abele, In Press) was also evident This suggests that comparisons of 6m (Rothera), 2-3m (South 367 Georgia) and inter-tidal (Signy) N. concinna will not be affected by population differences 368 between animals collected from different depths. N. concinna collected in the middle of summer 369 and tested within days of collection had no significant (t=0.57, p=0.57) differences in upper 370 lethal thermal limits (1°C per day) between those collected in the inter-tidal (11.6°C) or sub-371 tidal (6m, 10.8°C; SAM, unpublished data). The lack of any clear differences between Rothera

sub-tidal and Signy inter-tidal *N. concinna*, in the present study suggests that they respond
similarly to temperature, after a 2 month acclimation to 0.0°C.

374 Rothera N. concinna were also smaller than those from the other two locations, which 375 could introduce complicating factors due to growth and age differences as well as wider thermal 376 windows of small specimens (e.g. Pörtner 2002; Peck et al., 2007; Gsottbauer et al., 2007). 377 However, broad scale metabolic differences (see below), in the response to temperature, were 378 found between South Georgia and both Signy and Rothera N. concinna, which are unlikely to be 379 caused by size differences. Accepting these caveats, the present geographic comparisons within 380 the Southern Ocean, show for the first time specific responses of the adenylates, organic acids 381 and oxygen consumption to temperature, which suggest differences in critical limits and the 382 metabolic mechanisms employed to cope with acute temperatures close to these limits. There 383 was little evidence to suggest that Rothera or Signy N. concinna were temperature limited over 384 the range 0.0 to 10.0° C with measures of aerobic metabolism (oxygen consumption, citrate 385 synthase activity and ATP concentration) all increasing with temperature above 2.5°C. This 386 contrasts strongly with the pattern in L. elliptica which didn't show a reduction in these metrics 387 at temperatures above 2.5°C. This inicates they were suffering heat stress and have a narrower 388 thermal window than N. concinna.

389 In South Georgia N. concinna, citrate synthase activity and therefore the capacity of 390 aerobic metabolism in the mitochondria, increased with temperature in a similar fashion to 391 limpets from the other locations. However, in contrast to Rothera and Signy N. concinna, two 392 indicators of metabolism decreased in South Georgia N. concinna at higher temperatures. 393 Oxygen consumption peaked somewhere between 2.5 and 5.3°C and then declined whilst the 394 PLA pool, for buffering rapid ATP production, also declined above 5.1°C. Both of these suggest 395 a lower critical limit between 5.1 and 10.0°C (but perhaps as low as 2.5°C) for South Georgia N. 396 concinna, which was supported by the sharp increase in organic acids between 7.5 and 10.0°C 397 and therefore the recruitment of anaerobic pathways.

The wider thermal tolerance of *N. concinna* compared to *L. elliptica* is not surprising considering it ranges into the intertidal and can therefore be exposed to a more variable thermal environment. However, the difference in maximum littoral temperatures recorded so far, 12.3°C at Rothera (Waller et al., 2006) 10.7°C at Signy (Barnes et al., submitted) and 15.8°C at South Georgia (Davenport, 1997) was not reflected in any difference in the upper lethal limits for *N*.

403 *concinna* from different locations after 2 months acclimation to 0.0° C (10.0 to 12.5°C). 404 Davenport reported a short term survival limit of 15.6°C (12 hour exposure) for N. concinna at 405 South Georgia, higher than our medium term, 22 day, temperature limit of between 10.0 and 406 12.5°C. This is not unexpected as different thermal limits and protective mechanisms might be 407 expected if different rates of heating or cooling are used (Terblanche et al., 2007; Barnes et al., 408 2008; Peck et al., *in press*). This difference may also have been because N. concinna tested by 409 Davenport were acclimatised to summer conditions which could have raised their lethal limit 410 above that of animals acclimated to 0.0°C. However, it should also be noted that due to 411 behavioural adaptations, such as utilisation of cryptic habitats and evaporative cooling (Branch, 412 1981), body temperatures of limpets in the field are unlikely to reach these temperatures. The 413 maximum foot temperature of a Rothera N. concinna exposed on a sunny day has been shown to 414 be 8.8°C, although the average was much lower, 3.3 to 3.8°C, ranging from 1.5 to 9.1°C below 415 air temperature (Clark et al., 2008). If this relationship holds for South Georgia N. concinna then 416 foot temperature is unlikely to exceed their short-term (daily exposure) lethal limit during low 417 water emersion but might approach their medium term (weekly exposure) limit. Although N. 418 concinna are regularly exposed in the intertidal zone at Rothera and Signy, at South Georgia, it is 419 mainly a sub-tidal species (Davenport, 1997) and only rarely found above mean low water spring 420 tide level, reducing the risk of exposure to damaging high temperatures. The few inter-tidal 421 individuals found at South Georgia inhabit sheltered microhabitats and not exposed surfaces, 422 which are colonised by a siphonariid limpet, Kerguelenella lateralis. At its northern geographic 423 limit differences in habitat utilisation may be key to reducing the likelihood of *N. concinna* being 424 exposed to air temperatures close to its upper limit. They may therefore have evolved a high 425 anaerobic capacity to allow short-term, passive, survival during less frequent exposure (Pörtner, 426 2002). This fits with a growing body of evidence which shows that adaptation to life at range 427 limits may actually make animals less able to cope with further temperature increases (Stillman, 428 2003, Tomanek, 2005, Deutsch et al., 2008).

The passive survival strategy of *N. conccina* appears to be effective over several days, but is unlikely to be effective over longer time scales (Pörtner, 2006). After initial shock, animals may switch from acute mechanisms that allow short-term "passive" survival to longer-term acclimatory mechanisms with associated changes in tissue biochemistry and structure. The greater thermal flexibility of *N. concinna* than *L. elliptica* was further evidenced by the increase in adenylate concentration with temperature which suggests that there was an acclimatory
increase of the adenylates over the 2 week duration of the experiment, in an attempt to meet the
increase in energy demand with temperature.

437 Two month long incubations of *N. concinna* from South Georgia, Signy and Rothera, 438 long enough for seasonal acclimation, led to a reduction in the surface area of mitochondrial 439 cristae per unit muscle fibre volume (Morley et al., submitted). Latitudinal comparisons showed 440 that acclimated *N. concinna* from South Georgia had a higher citrate synthase activity than 441 Rothera N. concinna, which was negatively correlated with mitochondrial density across 442 locations. This strongly suggested an adaptation for fewer, more efficient mitochondria in 443 acclimated N. concinna from the northern limit of their range (Morley et al., submitted). Tissue 444 samples of *N. concinna* acclimated to 3.0°C are currently being analysed to investigate if 445 changes in aerobic pathways were accompanied by a change of anaerobic scope. These 446 differences in the mechanisms underlying, and the limits of, thermal tolerance between N. 447 concinna from South Georgia and both Signy and Rothera N. concinna are supported by genetic 448 differences, which suggest South Georgia N. concinna are a separate population (Beaumont and 449 Wei, 1991).

450 The Scotia arc region of the Southern Ocean has one of the fastest warming climates on 451 the planet (Meredith and King, 2005). If this rate of warming is faster than the rate of 452 evolutionary adaptation, then the acclimatory ability of individuals, populations and species is 453 likely to be a key determinate of their survival (e.g. Peck, 2008). Species with wider geographic 454 ranges are thought to be more resistant to extinction (Jablonski, 1986), which may in part be due 455 to the wider range of environments they experience. This should in turn lead to wider genetic 456 variation and therefore an increased physiological capacity to cope with environmental 457 perturbations (the climate variability hypothesis; Stevens, 1989). The current study has found 458 differences in thermal response of marine ectotherms between locations on the Scotia Arc and 459 Antarctic Peninsula. Investigations of a wider range of species will confirm if these differences 460 are an adaptive response to differences in experienced environment. Latitudinal comparisons 461 testing for differences in capacity between animals from different locations within the Southern 462 Ocean could be a key tool for predicting the vulnerability of species to the current trend of 463 warming.

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- 472

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 Sipunculus nudus below a critical temperature. J. Comp. Physiol. B 166, 492-500.
- 581
- 582 Table 1. Site details where limpets, *N. concinna* and the soft shelled clam *L. elliptica* were
- 583 collected during cruise JR109 and at Rothera research station.
- 584

Species	Site	Depth/m	Co-ordinates	Location
N. concinna	South Cove	6	67°34.25'S,	Rothera Point,
			68°08.00'W	Adelaide Island
	Borge Bay	0-1	60°42.16'S,	Signy, South Orkney
			45°35.45'W	Islands
	King Edward Point	2-3	54°17.03'S,	South Georgia
			36°39.30'W	
L. elliptica	Hangar Cove	10-20	67°33.92'S,	Rothera Point,
			68°07.67'W	Adelaide Island
	Borge Bay	10-20	60°42.16'S,	Signy, South Orkney
			45°35.45'W	Islands

586 Table 2. Shell lengths for limpets, *N. concinna* and the mud clam *L. elliptica* sampled for tissue

587	biochemistry. Mean ± sem	(number of individuals in	n brackets). With	in each species shel	l lengths

588	not sharing a common s	superscript are	significantly	different ($p < 0.05$).

Species	Location	Shell length/mm
N. concinna	Rothera	27.7±0.7 (20) ^a
	Signy	33.9±0.8 (20) ^b
	South Georgia	33.2±0.7 (20) ^b
L. elliptica	Rothera	74.0±3.9 (16) ^c
	Signy	84.2±3.7 (16) ^c

590

Table 3 The concentration of selected biochemical measures in *Laternula elliptica* foot muscle as temperature was increased in 5.0°C steps, after 1-week incubation at each temperature. Mean ±sem. Within each parameter, means not sharing a common superscript were significantly different (p<0.05).

Location	Parameter	Temperature			
		0.0	2.5	5.1	7.5
Rothera	O ₂ consumed	73.3±12.1 ^{ab}	83.4±16.4 ^a	$85.4{\pm}16.0^{ab}$	45.8±3.6 ^b
Signy	O ₂ consumed	49.6±4.8 ^{ab}	94.9±21.7 ^a	64.2±14.3 ^{ab}	34.2±2.6 ^b
Rothera	Heart rate	4.5±0.4	5.1	6.1	-
Signy	Heart rate	4.1±0.4	5.3±0.9	7.2	-
Rothera	CS activity	$0.57{\pm}0.05^{ab}$	0.65 ± 0.06^{a}	$0.40{\pm}0.14^{b}$	$0.44{\pm}0.10^{b}$
Signy	CS activity	0.38±0.07 ^{ab}	0.61 ± 0.05^{a}	$0.37{\pm}0.08^{b}$	0.30±0.02 ^b
Rothera	Arginine	3.0±1.2	3.1±0.4	1.5±0.2	8.3±1.2 ^a

Signy	Arginine	1.5±2.6	1.4±0.2	3.1±0.7	7.4±0.6 ^a
Rothera	AMP	0.08±0.01	0.07±0.005	0.03±0.006	$0.22{\pm}0.07^{a}$
Signy	AMP	0.04 ± 0.007	0.03 ± 0.003	0.07 ± 0.02	$0.53{\pm}0.22^{a}$
Rothera	ADP	0.60 ± 0.02	0.53 ± 0.02	0.41 ± 0.02	$0.90{\pm}0.2^{a}$
Signy	ADP	0.44 ± 0.03	0.42 ± 0.02	0.56±0.03	$0.79{\pm}0.07^{a}$
Rothera	ITP	0.11 ± 0.007^{a}	$0.10{\pm}0.00^{ac}$	$0.090 {\pm} 0.008^{ab}$	0.067 ± 0.006^{bc}
Signy	ITP	$0.06{\pm}0.01^{b}$	$0.08{\pm}0.004^{ab}$	0.10±0.005 ^{ac}	0.06 ± 0.007^{b}
Rothera	Acetate	0.0 ± 0.0	87.0±48	62.1±27.2	1182.7 ± 324^{a}
Signy	Acetate	0.0 ± 0.0	0.0±0.0	52.9±15	1251 ± 267^{a}
Rothera	Propionate	0.0 ± 0.0	0.0±0.0	0.0±0.0	$540.7{\pm}141^{a}$
Signy	Propionate	0.0 ± 0.0	0.0±0.0	0.0±0.0	586 ± 45^{a}
Rothera	pHi	7.6 ± 0.02^{a}	7.4 ± 0.01^{b}	$7.5{\pm}0.03^{b}$	$7.1 \pm 0.03^{\circ}$
Signy	pHi	7.6 ± 0.02^{a}	7.4 ± 0.02^{b}	7.5 ± 0.01^{b}	7.0 ± 0.09^{c}

598 Figure Legends

599 Fig. 1. The concentration of selected biochemical measures in *Laternula elliptica* foot muscle as

600 temperature was increased in 5.0°C steps, after 1-week incubation at each temperature; PLA

601 (phospho-L-arginine), ATP, cCO₂, (tissue carbon dioxide) and succinate. Filled bars, Rothera;

602 Open bars, Signy. Within each parameter, means not sharing a common superscript were

603 significantly different (p < 0.05).

604

Fig. 2. The concentration of adenylates in *Nacella concinna* foot muscle as temperature was

606 increased in 5.0°C steps, after 1-week incubation at each temperature. * indicates that AMP

607 concentrations for 10.0°C Rothera *N. concinna* were significantly different from all locations

608 except Signy *N. concinna* at 2.5°C and South Georgia *N. concinna* at 10.0°C (*p*<0.05). a

609 indicates that ADP levels were higher at 10.0°C than at other temperatures. Superscripts a and b

610 indicate that the PLA (phospho-L-arginine) concentration for South Georgia *N. concinna* was

611 lower at 10.0°C than either 0.0 or 5.1°C. Superscript c indicates that ATP concentration was

612 higher at 10.0 than 2.5° C. Mean \pm SEM.

613

Fig 3. The concentration of organic acids in *Nacella concinna* foot muscle as temperature was increased in 5.0°C steps, after 1 week incubation at each temperature. Points with superscript a are significantly different from all other treatments (p < 0.05). * indicates that the succinate concentration of South Georgia *N. concinna* at 10.0°C was significantly higher than all treatments except Rothera 7.5°C. Mean±SEM.

619

620 Fig. 4. The CS activity of foot muscle and the oxygen consumption of *Nacella concinna* as

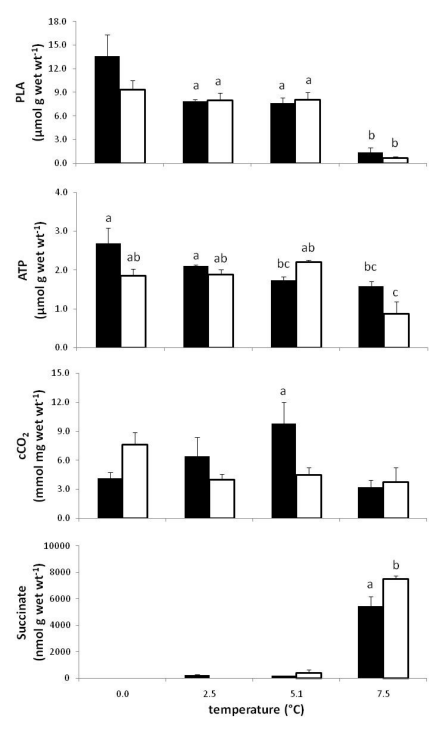
621 temperature were increased in 5.0°C steps, after 1 week incubation at each temperature.

622 Superscripts indicate that for each location CS activity was significantly higher at 7.5 and 10.0°C

than at lower temperatures (p < 0.05). The complex significant differences in oxygen consumption

between temperatures and locations are explained in the text. Mean \pm SEM.





627 Fig. 2

