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Exposure to solar radiation drives organismal vulnerability to climate: Evidence from an intertidal limpet



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ABSTRACT

Understanding the physiological abilities of organisms to cope with heat stress is critical for predictions of species' distributions in response to climate change. We investigated physiological responses (respiration and heart beat rate) of the ectotherm limpet *Patella vulgata* to heat stress events during emersion and the role of seasonal and microclimatic acclimatization for individual thermal tolerance limits. Individuals were collected from 5 microhabitats characterized by different exposure to solar radiation in the high intertidal zone of a semi-exposed rocky shore in winter and summer of 2014. Upper thermal tolerance limits (heat coma temperatures – HCTs, and heart rate Arrhenius break temperatures – ABTs) were determined for individuals from each microhabitat in both seasons under laboratory conditions. While we found a clear seasonal acclimatization, i.e., higher HCTs and ABTs in summer than in winter, we did not find evidence for microhabitat-specific responses that would suggest microclimatic acclimatization. However, operative limpet temperatures derived from in-situ temperature measurements suggest that individuals from sun exposed microhabitats have a much narrower thermal safety margins than those from less exposed surfaces or within crevices. Microhabitat specific thermal safety margins caused by high thermal heterogeneity at small spatial scales and the lack of short term acclimatization will likely shape small scale distribution patterns of intertidal species in response to the predicted increase in the frequency and intensity of heat waves.

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1. Introduction

Climate-related changes in species' distributions have already been observed in benthic systems (Birchenough et al., 2015). Predicting future species distributions and biodiversity patterns in a changing climate has become a priority over the last decades (Borja, 2014) in order to anticipate the ecological and socio-economic impacts of climate related changes on local biodiversity (Sarà et al., 2014). Therefore, knowledge on the physiological abilities of ectotherms in response to changes in temperature is critical (Bozinovic et al., 2011). Organismal physiological

performance and fitness affect population dynamics, species interaction and ecosystem functioning and are driven by the local environmental context (Doney et al., 2012). For instance, intertidal invertebrates face high spatial and temporal temperature heterogeneities caused by exposure to water and air over the tidal cycle, and microhabitat-specific exposure to solar radiation in different seasons (Chapperon and Seuront, 2011; Seabra et al., 2011). The microtopography of rocky substrates strongly influences the degree of exposure to solar radiation and shapes the variability in substrate temperature observed over the tidal cycle and in different seasons (Harley, 2008; Miller et al., 2009; Seabra et al., 2011).

Here we investigate whether microhabitat- and season-specific thermal histories of the limpet *Patella vulgata* affect its physiological ability to cope with sublethal thermal stress during emersion. Specifically, we examine the physiological responses of *P. vulgata* (respiration and cardiac activity) to heat stress in the laboratory and derived thermal safety margins for five microhabitats

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Table 1
Definitions of thermal physiological descriptors used in the present study.

Thermal physiological descriptor	Definition
Heat Coma Temperature (<i>HCT</i>)	Temperature at which individuals became detached from support and did not show any response after pricking the foot and tentacles with a needle (Evans, 1948). Individuals showed a response to a test of irritability after 24 h. <i>HCT</i> was used as a proxy for the critical thermal limit (CT_{max}) of locomotor performance (Polgar et al., 2015)
Arrhenius Break Temperature (<i>ABT</i>)	Temperature above which heart beat rates decrease rapidly (Stillman and Somero, 1996), thus indicating cardiac function failure.
Thermal safety margin (<i>TSM</i>)	Difference between the upper thermal tolerance limit (defined as <i>HCT</i> in the present study) and the maximal estimated operative temperature (Sunday et al., 2014) recorded over a 14 days period in summer 2014 within each studied microhabitat.

characterized by contrasting exposure to solar radiation. We hypothesize that both microhabitat (short-term) and seasonal (long-term) thermal acclimatization in *P. vulgata* affect its thermal tolerance limits and physiological performance in response to heat stress. In particular, we expect individuals from microhabitats characterized by large daily temperature variations to show a greater ability to cope with thermal stress than individuals from microhabitats with more stable thermal conditions (Magozzi and Calosi, 2014). Similarly, we anticipate seasonal variations in the thermal sensitivities and thermal tolerance limits of limpets (i.e., we expect summer-adapted individuals to show a greater capacity to cope with heat stress than winter-adapted individuals, Davies, 1965).

2. Materials and methods

2.1. Sampling site

Individuals of *P. vulgata* were collected from a semi-exposed rocky shore (Dellec, Plouzané, France, 48°21'09"N, 4°34'01"W) in winter and summer 2014 during daytime ebbing low tide. At the study site, *P. vulgata* occurred at densities of 135 ± 18 individuals/m². Since one of the objectives was to examine the effect of the recent thermal history on physiological responses to heat stress during emersion, individuals were collected one day before laboratory measurements to avoid acclimation to laboratory conditions. Adult *P. vulgata* belonging to a similar size class (with average height and length of ca. 15 mm and 32 mm) were collected at the upper limit of their intertidal distribution where they experience the greatest temperature variations (Seabra et al., 2011). Limpets were collected from five microhabitats characterized by different levels of exposure to solar radiation and thus potential differences in thermal stress: crevices (C, very low exposure), vertical north exposed (NE, low exposure), vertical west exposed (WE, high exposure), flat rock (FR, higher exposure) and vertical south exposed surfaces (SE, highest exposure).

Intertidal ectotherms' body temperatures are not correlated to the air temperature as they are affected by multiple interactive abiotic factors such as exposure to solar radiation, wind chill and wave splash (Helmuth and Hofmann, 2001; Fitzhenry et al., 2004; Broitman et al., 2009). Body temperatures also depend on morphological properties like shell shape and color. To account for the influence of this suite of environmental and morphological factors on operative body temperatures, we deployed biomimetic loggers (robolimpets, Lima and Wethey, 2009) adjacent to substrate temperature loggers (iButton thermochron DS1921 G-F5, Homechip UK). Temperature loggers were significantly correlated to robolimpets across all microhabitats ($R^2 > 0.95$, $p < 0.05$; see Fig. 1 in Supplementary materials as an example) and had an average bias of 1.25 °C and a mean square error of 1.33 °C. Due to logistic reasons, robolimpets were not deployed in all

microhabitats and seasons. Instead, DS1922L iButton loggers were attached to the rocky substrate in each microhabitat using glazing tapes (Hodgson Sealants, UK) and temperature measurements were taken every 10 min over 14 days in both seasons. Note that summer recordings were conducted in September 2014 which was the month with the highest average daily maximum temperature for the period 2011–2014 as evidenced by data collected using biomimetic loggers at a shore less than 50 km away from our study site (see Fig. 2 in Supplementary materials). Operative limpets temperatures were thus estimated using the equations of the significant linear relationships obtained between robolimpets and iButtons temperature in the different microhabitats (estimated operative limpet temperature hereafter).

2.2. Laboratory experiments

We assessed three descriptors of thermal tolerance during emersion: heat coma temperature, Arrhenius break temperature and thermal safety margin (*HCT*, *ABT* and *TSM*, respectively; see Table 1 for definitions).

Upon collection, individuals were brought to the laboratory within 30 min and placed into a tank with natural tidal cycle and 12 h photoperiod. Shells were scraped and carefully cleaned with ethanol to remove epibionts. To assess *HCT*, individuals were directly placed on the vertical walls of two cubic transparent plastic tanks (13 cm side). For combined respiration and heart beat rate measurements or heart beat rate measurements only, limpets were positioned on horizontal sandstone plates after gluing heart beat infrared sensors onto their shells. Animals were sprayed with seawater to allow replenishment of mantle water and attachment to their support (Williams et al., 2005). Seawater temperature was similar to that experienced by individuals at the collection site during immersion (10 °C in winter and 16 °C in summer). Room temperature was maintained to that of the seawater.

In all experiments, thermal stress during emersion was simulated by immersing containers (tanks or respirometry bottles without any water depending on experiments) within a thermostated water bath (Huber, Cryo-Polystat CC –20/200 °C 12 L, Avanteq, France). Two similar thermal ramps were used in this study, both featuring a warming rate of 6 °C/h. However, for respiration and cardiac rate measurements, the water bath temperature was instantaneously raised 6 °C at the beginning of each hour, and kept constant until the next hour. On the other hand, on heat coma experiments and heart rate measurements, water bath temperature was raised using a smoother ramp of 1 °C every 10 min (i.e., each target temperature was held for ca. 10 min). Data loggers (DS1921G-F5 iButton thermochron, Homechip UK) were placed inside each tank (three per tank) and each bottle (one per bottle) to confirm that water bath temperature and the temperature inside each tank were similar. An absorbent tissue saturated with seawater was placed in each tank/bottle to maintain 100% humidity level.

2.2.1. Upper critical thermal limit measurement

Upper thermal tolerance limits were determined by assessing HCTs. In this study, thermal coma was defined sensu stricto as the absence of physical response following stimuli – i.e. by pricking the foot and tentacles with a needle (Evans, 1948). HCTs were determined by using the smoother temperature ramp (1 °C/10 min) and registering the temperature at which animals started to detach from the container walls and failed to retract their foot and tentacles when gently stimulated with a needle (Clarke et al., 2000a, 2000b). After reaching HCTs, individuals were placed inside the tidal tank at field temperature and their survival was checked after 24 h. Individuals detached from the substratum and not responding to any stimuli were considered dead. Since we were interested in heat coma temperatures and not lethal temperatures, data from dead individuals were discarded from the analyses. HCTs were estimated for individuals from each microhabitat in both seasons as the arithmetic mean of the temperatures at which individuals reached the endpoint (n=22 per microhabitat, experiments repeated twice in each season).

2.2.2. Respiration and cardiac activity measurements

Simultaneous monitoring of cardiac activity and respiration were used to measure metabolic responses of limpets to heat stress. Heart beat frequencies were measured with infrared sensors that enable non-invasive measurements of heart beats through the shells (Burnett et al., 2013). On the day of each experiment, individuals were equipped with IR sensors (n=5; one individual per microhabitat) and moved into airtight 500 ml plastic bottles adapted to allow the simultaneous measurements of respiration (CO₂ production) and cardiac activity. Experiments were repeated four times in winter and five times in summer. Plastic bottles were immersed in the thermostated water bath at field water temperature (10 °C in winter, 16 °C in summer). Individuals were moved with their sandstone plates into bottles to minimize handling stress before the start of the experiment. As explained above, water bath temperature was instantaneously increased 6 °C at the beginning of each hour from 10 °C (winter) or 16 °C (summer) until a target temperature of 43 °C was reached. Respiration measurements were taken in each bottle successively after 20 min of acclimation to the target temperature. Bottles were connected to a closed circulation system with an integrated infrared CO₂ analyser (Li-Cor, Li-820) and a desiccation column filled with anhydrous calcium sulphate (Drierite, Xenia, USA). An adjustable pump maintained air flow at 0.8–0.9 L min⁻¹. CO₂ partial pressure (pCO₂, in ppm: parts per million) in each bottle was recorded using the Li-820 software. pCO₂ was measured every 5 s over a 3 min period (Tagliarolo et al., 2013). Respiration rates within each bottle were estimated by determining the linear slope of pCO₂ over time (see Fig. 3 in Supplementary materials). Cardiac activity was recorded every 10 min for 60 s during the whole thermal ramping essay. At each temperature along the thermal ramp (10 °C and/or 16 °C, 22 °C, 28 °C, 34 °C, 40 °C, 43 °C), variations in pCO₂ were measured within each bottle successively, with a pause of several minutes between measurements to stabilize the system. An entire cycle of respiration measurements lasted ca. 30 min. Preliminary measurements were conducted with empty bottles over the same thermal ramp and pCO₂ variations were negligible in comparison to pCO₂ increases in the presence of individual limpets. Fluxes of pCO₂ were corrected for the net volume of the system and incubation time and normalized to biomass (μmol CO₂ gAFDW⁻¹ h⁻¹). Individual biomass was estimated as ash-free dry weight (AFDW) through loss on ignition (60 °C for 48 h for drying and 450 °C for 4 h for burning).

2.2.3. Arrhenius breakpoint measurements for cardiac activity

ABT was calculated for each microhabitat/season combination. Individuals fitted with heart beat sensors were exposed to a

step-wise ramp of 1 °C every 10 min, from 10 °C (winter) or 16 °C (summer) up to 43 °C or until heart beat was lost). Animals were kept on sandstone plates inside cubic plastic tanks (13 cm side) partially submerged in a water bath. ABTs were calculated as the temperature corresponding to the intersection point of the two best fitting lines on each side of the Arrhenius plots (Stillman and Somero, 1996).

2.3. Statistical analyses

The distributions of thermal physiological descriptors (Table 1) at both seasons and in the different microhabitats were checked for normality using the Kolmogorov–Smirnov test. Levene's test was used to test for equality of variances. When data were normally distributed, a two-factor analysis of variance was used to determine the effects of season, and microhabitat on respiration, heart beat rate and Arrhenius break temperature. The effects of temperature upon both respiration and cardiac rates were assessed using repeated measures ANOVA with a Greenhouse-Geisser Correction since the data violated the assumption of sphericity (Mauchly's test of sphericity). Post hoc tests incorporating the Bonferroni correction were done to distinguish differences in respiration and heart rates between temperatures. When data failed normality even after log transformation, the non-parametric Mann-Whitney *U* test (MW test hereafter) was used to compare heat coma temperatures between seasons. Similarly, the Kruskal Wallis Test (KW test hereafter) followed by subsequent non-parametric post hoc analyses (based on the Tukey test; Zar, 2010) were used to compare heat coma temperatures and thermal safety margins between individuals collected within different microhabitats. All statistical analyses were carried out using IBM SPSS Statistics 20 (IBM Corp. Released 2011).

3. Results

3.1. Estimated operative temperatures of *Patella vulgata*

From the 10 deployed temperature data loggers (2 per microhabitat), 8 and 7 provided reliable data in winter and summer, respectively. In both seasons, minimal variations in estimated operative limpet temperatures were observed in microhabitats least exposed to solar radiation, whilst maximal thermal variations occurred on the sun exposed surfaces (Table 2). In addition, during diurnal low tides, maximum estimated operative limpet temperatures and temperature variability were obtained on surfaces directly exposed to solar radiations (Figs. 1 and 2). In contrast, estimated operative limpet temperatures within crevices and on north exposed rocks were much less variable in both seasons (Figs. 1 and 2).

3.2. Seasonal effect upon *Patella vulgata* thermal sensitivity and limits

In both seasons, respiration and cardiac rates were significantly affected by temperature (Table 3). In winter, respiration rates increased from 16 °C to 28 °C and then stabilized until the end of the thermal ramp (Table 3, Fig. 3). In winter, cardiac activity significantly increased from 10 °C to peak at 34 °C and then slowed down to almost acardia at 43 °C (Fig. 3). In summer, respiration rates were significantly lower at 16 °C than any other temperature (except 22 °C). Respiration rates were mostly stable over the range 22–43 °C (but see significant differences in Table 3). Heart beat rate increased from 16 °C to 34 °C (Fig. 3) but no significant difference in heart beat rate was registered over the range 34–43 °C (Table 3).

Table 2

Patella vulgata estimated operative temperature in winter and in summer 2014 on a semi exposed rocky shore (Dellec, 48°21'09N, 4°34'01W, France) in different microhabitats (C: crevice; NE: vertical north exposed; WE: vertical west exposed; FR: flat rock; SE: vertical south exposed). n=1990 and n=1988 in all microhabitats in winter and summer respectively (except n=702 in summer on vertical west-exposed surfaces).

Season	Microhabitat	Mean (\pm SE)	Min–Max	Range
Winter	C	8.2 (0.0)	3.3–13.2	9.9
	NE1	8.9 (0.0)	5.5–12.3	6.8
	NE2	8.6 (0.0)	5.5–11.9	6.4
	WE	8.5 (0.0)	4.2–13.7	9.5
	FR1	8.2 (0.0)	3.7–15.2	11.4
	FR2	8.1 (0.0)	3.3–13.7	10.5
	SE1	8.4 (0.0)	4.2–18	13.8
	SE2	8.5 (0.0)	4.7–19.5	14.8
Summer	C	18.2 (0.0)	15.7–21	5.3
	NE	15.2 (0.0)	13.1–26.3	13.2
	WE	17.0 (0.1)	14.2–29.5	15.2
	FR1	18.1 (0.1)	13.3–34.7	21.4
	FR2	18.0 (0.1)	13.7–34.2	20.5
	SE1	19.3 (0.1)	14.3–40.4	26.2
	SE2	20.3 (0.2)	14.2–39.0	24.8

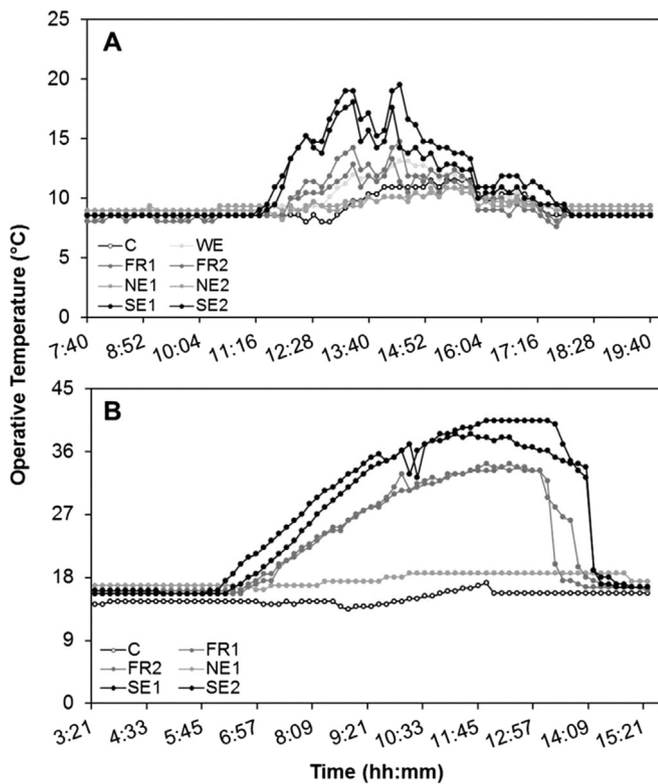


Fig. 1. *Patella vulgata* estimated operative temperature during two successive high tides in winter (A) and in summer 2014 (B) on a semi exposed rocky shore (Dellec, 48°21'09N, 4°34'01W, France) in different microhabitats (C: crevice, white dots; WE: vertical west exposed, light grey dots; NE: vertical north exposed, light grey dots; FR: flat rock, dark grey dots; SE: vertical south exposed, black dots). Note that no temperature was recorded on WE in summer on that specific low tide.

The two-factorial analysis of variance revealed a significant seasonal effect on (i) heart beat rate ($p < 0.01$, Table 4) at different temperature along the thermal ramp (but not on respiration rate, $p > 0.05$, Table 5) and also on (ii) ABT values ($F=19.373$, $df = 1$, $p < 0.001$). More specifically, heart rate was higher in winter than in summer at all temperatures between 16 °C and 34 °C, and at 43 °C ($p < 0.05$). ABTs were higher in summer than in winter (Table 6, Fig. 4). Similarly, heat coma temperatures (HCTs) were

significantly higher in summer (42.5 ± 0.1 °C) than in winter (38.2 ± 0.1 °C; MW test, $Z = -16.067$, $p < 0.001$; Fig. 5).

3.3. Microhabitat effect upon *Patella vulgata* thermal sensitivity and limits

The two-factorial analysis of variance revealed no significant effect of microhabitat upon respiration and cardiac rates along the thermal ramp (Tables 4 and 5), and upon ABTs ($F=0.390$, $df = 4$, $p=0.390$).

HCTs were significantly higher in organisms collected on flat rocks than in individuals collected in any other microhabitat in winter (Tukey test following KW Test, $p < 0.05$, Fig. 5). In summer, the temperature at which heat coma was induced was significantly higher in individuals from vertical north- and south-faced surfaces than in animals from crevices and flat rock surfaces ($p > 0.05$, Fig. 5).

Thermal safety margins were significantly different between microhabitats (KW test, $\chi^2=186.231$, $df = 4$, $p < 0.01$, Fig. 6). On average, TSMs were significantly larger on individuals collected within crevices than on individuals from any other microhabitat (Fig. 6). TSMs of limpets from vertical west- and north-exposed microhabitats were significantly higher than those from flat and south exposed rocks (Fig. 6). The greatest TSM (22.5 °C) was observed in an individual collected within a crevice whilst the smallest TSM was recorded on vertically south-faced substrate (0.6 °C).

4. Discussion

4.1. Sun exposure as a driver of microscale heterogeneity in *P. Vulgata* operative temperatures

Patella vulgata experiences largely different thermal variations and extremes in different microhabitats, specifically during daytime low tides. Individuals from sun exposed microhabitats like flat, or vertical west- and vertical south- exposed rocks face variations in body temperature of up to 20 °C whilst emerged in summer (Figs. 1 and 2). As shown for the limpet *Lottia gigantea*, intertidal organisms in sun exposed microhabitats frequently experience sublethal and sometimes lethal thermal stress (Miller et al., 2009). Crevices and north-faced surfaces appear to be more stable microenvironments with much lower thermal fluctuations and maxima. This study thus reinforces the evidence that the amount of solar radiation received by intertidal ectotherms within microhabitats drives the mosaic patterns of body temperatures and thermal stress at the microscale (Marshall et al., 2010; Lima et al., 2015). The effect of sun exposure is such that body temperature variations between sun exposed and shaded microhabitats have been demonstrated to be greater than those observed between shore levels or between sites separated by hundreds of km (Seabra et al., 2011).

Besides solar radiation exposure, other local factors such as wave splash and timing of low tides have been shown to drive counterintuitive patterns of invertebrate body temperatures (Helmuth et al., 2006) and thermal tolerance may vary over the distribution range of a given species (Kuo and Sanford, 2009). Intra-specific variability in physiological and biochemical adaptations of marine ectotherms to thermal stress has also been documented for different shore levels according to the emersion gradient (Stirling, 1982; Tomanek and Somero, 1999; Stillman and Somero, 1996; Davenport and Davenport, 2005; Stenseng et al., 2005; Nguyen et al., 2011; Prusina et al., 2014). Similarly, previous studies have highlighted differences in the metabolic responses to heat stress of intertidal ectotherms from different microhabitats.

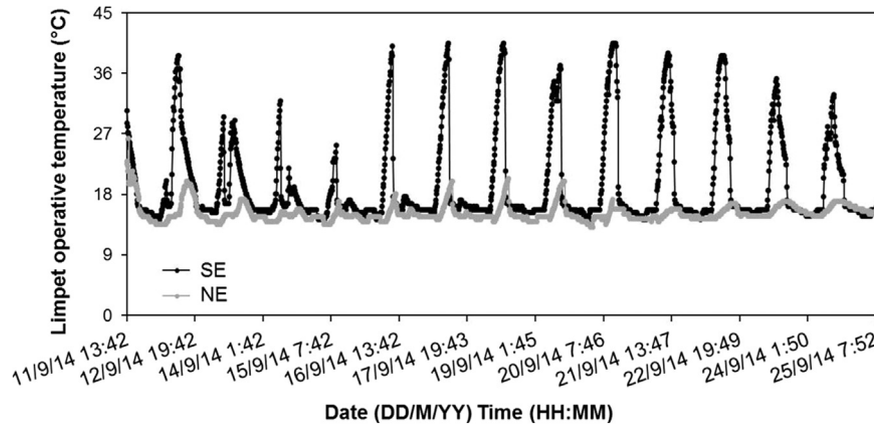


Fig. 2. *Patella vulgata* estimated operative temperature values recorded over 14 days every 10 min in summer 2014 on a semi exposed rocky shore (Dellec, 48°21'09N, 4°34'01W, France) on south- and north- exposed surfaces (black and grey dots and lines, respectively).

For instance, the limpet *Cellana tramoserica* displays higher metabolic rate (oxygen consumption) in response to heat stress in stable environments (rock overhangs) than in variable microhabitats (exposed flat rock, Sinclair et al., 2006).

4.2. Lack of short term acclimatization in *P. Vulgata* thermal sensitivities and tolerance

In this study, individuals did not show any difference in their respiratory or cardiac responses to heat stress, regardless of the extent of daily thermal fluctuations and the frequency of exposure to thermal extremes that the limpets had experienced within their microhabitats. We found some significant differences in the upper thermal limit (HCTs) between *P. vulgata* from different microhabitats, but no clear pattern was detected that would suggest that the acclimatization to the microhabitat specific thermal conditions affects the limpets' ability to cope with heat stress. The lack of microhabitat specific response was confirmed by ABTs for heart rates that did not significantly differ among individuals from different microhabitats. These results seem counterintuitive since reversible acclimatization should be an advantage in individuals experiencing diel temperature variations (Gabriel, 2005; Young and Gifford, 2013). Nonetheless, others species such as the high shore snail *Echinolittorina malaccana* also show no significant differences in metabolic rate between sun-exposed and shaded microhabitats when facing heat stress (Marshall et al., 2013). Besides the lack of short term acclimatization to surrounding microclimatic conditions, *P. vulgata*, as others ectothermic species, adapts its metabolism through thermal compensation (e.g. Marshall et al., 2013). Even though the high coefficient of variations in respiratory rate (25–67% in winter, and 43–63% in summer) is indicative of a high inter-individual variability, on average, respiration did not vary significantly in the range 22/28–43 °C. This temperature-insensitive metabolism has previously been observed in terrestrial ectotherms like reptiles (Seebacher, 2005) and marine invertebrates like the snails *Littorina littorea* during the whole

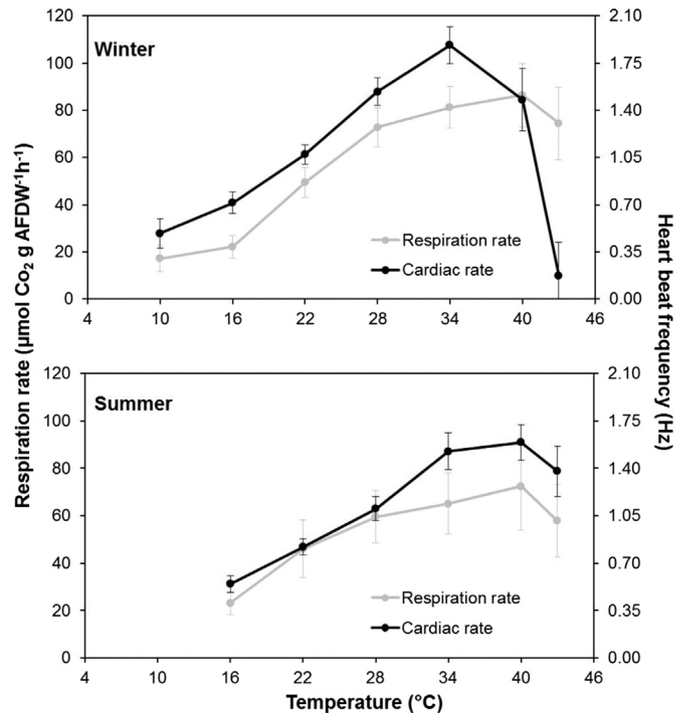


Fig. 3. *Patella vulgata* respiration rates (grey dots) and cardiac activity (black dots) in response to temperature in winter and in summer. Values are means and error bars are 95% confidence limits. 17 <math>\leq \eta_{\text{respiration-winter}} < 21</math>, 12 <math>\leq \eta_{\text{respiration-summer}} < 21</math>, 20 <math>\leq \eta_{\text{cardiac activity-winter}} < 21</math>, 14 <math>\leq \eta_{\text{cardiac activity-summer}} < 19</math>.

resting state (Newell and Pye, 1970, 1971), *E. malaccana* (Marshall et al., 2013) over a temperature range of 35–46 °C, and the limpet *Lottia digitalis* over a temperature range of 15–35 °C (Bjelde and Todgham, 2013). Others *Lottia* limpets (*L. scabra*, *L. austrodigitalis*, *L. limatula*, *L. pelta*) increase their aerial respiration rate with

Table 3
Repeated measures ANOVA with a Greenhouse-Geisser Correction followed by Post hoc tests using the Bonferroni correction to investigate differences in the respiratory and cardiac responses of *Patella vulgata* at different temperatures over a thermal ramp in the laboratory (increase in 6 °C increments) in both winter and summer.

Season	Parameter	dfs	F	P	Post Hoc
Winter	Respiration	2.370, 37,919	55.967	< 0.001	10=16 < 22 < 28 = 34 = 40 = 43; 22=43
	Heart rate	2.542, 45.757	69.888	< 0.001	10 < 16 < 22 < 28 < 34 = 40 > 43; 10 = 43; 22 = 40; 28 = 40
Summer	Respiration	2.246, 24.709	14.983	< 0.001	16 = 22 = 28 = 34 = 40 = 43; 16 < 28; 16 < 34; 16 < 40; 16 < 43; 22 < 40; 34 > 43
	Heart rate	2.495, 32.431	41.626	< 0.001	16 < 22 < 28 < 34 = 40 = 43; 28 = 43

Table 4

Two-factorial analysis of variance to investigate season, microhabitat and season × microhabitat effects upon cardiac rates of *Patella vulgata* at different temperatures along a thermal ramp in laboratory (increase in 6 °C increments).

Cardiac rate	15 °C			22 °C			28 °C			34 °C			40 °C			43 °C		
	df	F	P	df	F	P	df	F	P	df	F	P	df	F	P	df	F	P
Microhabitat	4	1.445	0.244	4	1.418	0.253	4	1.761	0.163	4	0.200	0.937	4	0.697	0.600	4	2.005	0.117
Season	1	8.918	<0.01	1	25.527	<0.001	1	37.806	<0.001	1	10.730	<0.01	1	0.897	0.352	1	14.052	<0.01
Microhabitat × microhabi	4	2.082	0.109	4	2.141	0.101	4	2.637	0.053	4	0.223	0.923	4	0.543	0.705	4	1.759	0.161

increasing temperature up to a temperature threshold beyond which respiration decreases (Miller et al., 2015). Nevertheless, these species also show some metabolic rate control over the range 20–30 °C (Q_{10} values lower than 1.5, Miller et al., 2015). Although short-term metabolic compensation (maintenance of constant rate processes) is not universal (e.g. Bannister, 1974), it is similarly to metabolism depression (decrease of metabolic rate below that of resting state), an adaptation strategy under environmental (thermal) change. Both temperature-independent metabolism and hypometabolism allow ectotherms to minimize energy loss (energy conservation) and withstand higher temperatures for longer periods during aestivation (Guppy and Withers, 1999; Marshall et al., 2011). The lack of shifts in thermal sensitivity between individuals from different microhabitats suggests that *P. vulgata* is a generalist species, metabolically adapted to the wide range of temperatures that it can potentially encounter in the different microhabitats on the upper intertidal zone. However, being generalist also means that individuals living on sun exposed surfaces are not better adapted to cope with the thermal extremes frequently reached in these microhabitats, as indicated by the lack of differences in their thermal tolerance parameters (HCTs and ABTs). This is not consistent with previous studies that indicated that the recent thermal history of individuals is the most influential factor determining thermal tolerance (Lutterschmidt and Hutchison, 1997; Clarke et al., 2000a). Even though the temperature experienced in the field during the studied period never surpassed that of the HCTs obtained in laboratory, some south exposed individuals had a very narrow thermal safety margin (0.6 °C). Limpets on south faced surfaces seem particularly vulnerable since the maximal temperature experienced in the field in summer during the study period (40.4 °C) exceeded that of the ABTs calculated based on our laboratory results. Other ectothermic species have shown an absence of microhabitat acclimatization regarding their upper thermal limits. Instead they displayed different responses to heat stress at the molecular and cellular levels. For instance, despite the lack of local adaptation in a physiological parameters (critical thermal maximum, CT_{max}), coastal crabs exhibited a higher production of heat shock proteins (Hsp70) in comparison with estuarine crabs (Madeira et al., 2012). These differences were associated with differences in local temperatures and air humidity (Madeira et al., 2012). Similarly, shrimp in the upper intertidal showed higher production of Hsp70 and total ubiquitin and histopathological alterations than shrimp from the

Table 5

Two-factorial analysis of variance to investigate season, microhabitat and season × microhabitat effects upon respiration rates of *Patella vulgata* at different temperatures over a thermal ramp in laboratory (increase in 6 °C increments).

Respiration rate	15 °C			22 °C			28 °C			34 °C			40 °C			43 °C		
	df	F	P															
Microhabitat	4	1.537	0.216	4	0.320	0.862	4	0.229	0.920	4	1.738	0.166	4	0.987	0.429	4	0.345	0.844
Season	1	0.003	0.957	1	0.136	0.715	1	2.242	0.144	1	3.376	0.075	1	1.526	0.226	1	1.412	0.247
Microhabitat × season	4	0.236	0.916	4	0.279	0.889	4	2.703	<0.05	4	0.907	0.471	4	1.172	0.342	4	1.549	0.221

Table 6

Arrhenius Break Temperatures of *Patella vulgata* heart rate obtained in individuals facing using a step-wise ramp of 1 °C every 10 min in laboratory in both winter and summer. Individuals were collected within different microhabitats (C: crevice; WE: vertical west exposed; NE: vertical north exposed; FR: flat rock; SE: vertical south exposed).

Season	Microhabitat	N	Mean ABT (± SE)	Min–Max	Range
Winter	C	5	35.3 (1.5)	30.3–38.8	8.5
	NE	6	33.3 (1.3)	28.4–37.5	9.1
	WE	4	36.1 (0.6)	35.1–37.5	2.4
	FR	7	34.2 (1)	31.1–36.7	5.6
	SE	5	34.4 (0.8)	32.4–37.0	4.6
Summer	C	11	36.2 (0.7)	30.6–38.8	8.2
	NE	9	37.6 (0.5)	35.3–40.6	5.3
	WE	6	36.8 (0.6)	34.8–38.4	3.1
	FR	6	37.3 (0.7)	35.7–40.3	4.6
	SE	7	37.9 (0.5)	35.9–40.0	4.1

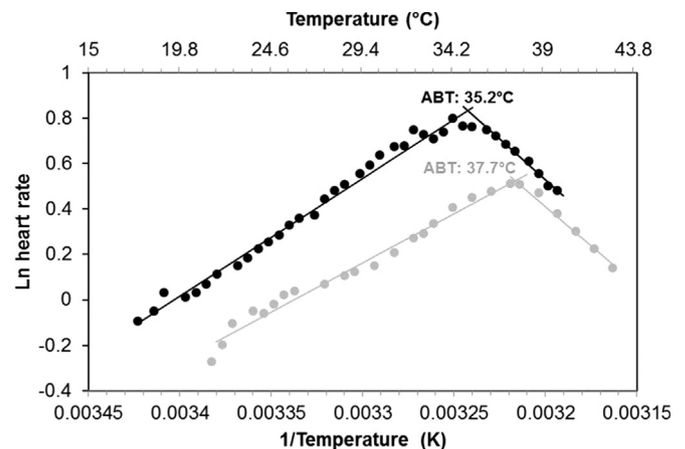


Fig. 4. Representative examples of Arrhenius plots used to define the Arrhenius Break Temperature (ABT) in *Patella vulgata* individuals in response to a step-wise ramp of 1 °C every 10 min in laboratory in winter (black dots and line) and in summer (grey dots and line).

lower intertidal despite an absence of differences in CT_{max} (Madeira et al., 2015). These studies stress the need to further investigate the heat stress response of *P. vulgata* at the molecular level, as the patterns of variations of stress proteins have been

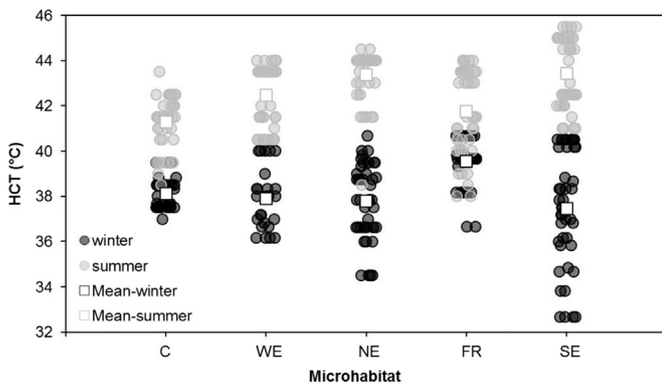


Fig. 5. Temperatures inducing heat coma (*HCTs*) in *Patella vulgata* individuals in response to a step-wise ramp of 1 °C every 10 min in laboratory in winter (black points) and in summer (grey points). Individuals were collected within different microhabitats (C: crevice; NE: vertical north exposed; WE: vertical west exposed; FR: flat rock; SE: vertical south exposed). Each dot represents a *HCT* value ($25 < n < 44$ in winter; $38 < n < 43$ in summer). Squares indicate the mean *HCT* obtained in individuals sampled within each microhabitat in both winter and summer.

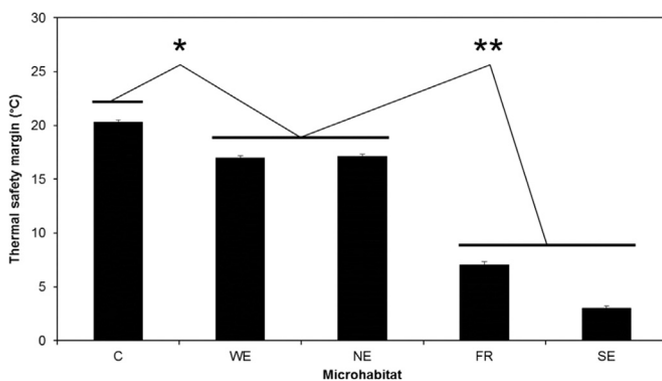


Fig. 6. Thermal safety margins determined for *Patella vulgata* individuals in different microhabitats (C: crevice; NE: vertical north exposed; WE: vertical west exposed; FR: flat rock; SE: vertical south exposed). TSM was defined as the difference between the *HCT* measured for each tested individual from a microhabitat and the highest estimated operative temperature recorded within that microhabitat over a 14 day period in summer 2014. Values are means (\pm SE, $38 < n < 43$). *: $P < 0.05$, **: $P < 0.01$.

shown to occur even within small-scale thermal mosaics (Hofmann, 1999).

4.3. Seasonal acclimatization in *P. vulgata* thermal sensitivities and tolerance

Even though the daily temperature variations at our study site were large, perhaps the unpredictability of the frequency of occurrence of heat stress events could have led to the absence of short term acclimatization of *HCTs* and *ABTs* (Clarke et al., 2000a). Indeed, even if acclimatization can occur over a few weeks (mitochondrial function) or days (cardiovascular function), the energy spent to increase *ABTs* and *HCTs* to maintain organismal homeostasis under stochastic events such as heat waves is potentially too high, outweighing its long term benefits. Another explanation is that diel variations in temperature at the study site were not high enough in comparison with long term thermal variations. This hypothesis is compatible with our results that clearly indicate seasonal acclimatization of heart beat rate i.e. cardiac activity was lower in winter than in summer at all temperatures (except at 40 °C) and heart beat rate peaked at a lower temperature in winter than in summer. Similarly, the greater *ABTs* and *HCTs* in summer are indicative of seasonal acclimatization of the thermal tolerance

for cardiac function and upper limit. Evidence of seasonal acclimatization in cardiac responses to heat stress and thermal tolerance is common in the literature. For instance, the limpet *Lottia limatula* (Segal, 1956), the pond snail *Lymnaea stagnalis* (Harrison, 1977) and the crab *Pachygrapsus marmoratus* (De Pirro et al., 1999) all show greater heart rate in winter than in summer at corresponding temperatures. The temperature for peak heart rate in *L. stagnalis* (Harrison, 1977) was also found to be lower in winter (30.5 °C) than in summer (37.5 °C). Seasonal acclimatization of the cardiac function allows individuals to be more tolerant to higher temperatures in the warm months.

As with other ectotherms such as the freshwater mussel *Unio tumidus*, there was no seasonal acclimatization in respiratory rates (Lurman et al., 2014), but we previously discussed an alternative strategy used by *P. vulgata* to cope with thermal stress, i.e. thermal insensitive metabolism. In summer, the higher upper thermal tolerance limits (*HCTs*, 42.5 ± 0.1 °C as compared to 38.2 ± 0.1 °C in winter) and thermal tolerance limits of heart function (*ABTs*, 36.9 ± 0.3 °C instead of 34.3 ± 0.6 °C in winter) most certainly facilitate *P. vulgata*'s ability to cope with temperature extremes and higher sublethal thermal stress experienced in the upper intertidal zone. Although this intra-individual flexibility in shifting upper thermal limits over extended periods of time may constitute an adaptive strategy in a warming climate, this study also reveals that not all individuals within the same shore level are equal in terms of vulnerability to future warming.

4.4. Not all vulnerable to the warming climate: the importance of microscale heterogeneity

High shore ectotherms are generally assumed to be particularly vulnerable to the predicted increase in the frequency and intensity of heat events (Planton et al., 2008) as they often live at or close to their upper thermal tolerance limits (Somero, 2002). However, our results highlight that not all high shore limpets are threatened by further warming and heat events. Since individuals of *P. vulgata* do not appear to show any acclimatization ability in their thermal tolerance and sensitivity as a function of their surrounding microclimatic conditions, those living on south-exposed and flat rocks seem to be highly vulnerable to extreme heat events with current *TSMs* (based on our 14 day-long thermal recordings) ranging from 3.3 °C to 9.3 °C, and 0.6–5 °C, respectively. At the opposite, organisms on vertical north- and vertical west-exposed rocky walls display a *TSM* of ca. 17 °C. *TSM* is even higher within crevices, exceeding 20 °C. This is consistent with a previous study that highlighted differences in safety margin between shade-resting and sun exposed *E. malacana* snails (Marshall et al., 2013). The tropical freshwater snail *Clea nigricans* also showed much higher thermal safety margin than expected (Polgar et al., 2015). These results show the need to investigate the impact of thermal heterogeneity on the patterns of thermal tolerance within populations at the microscale.

5. Conclusion

This study emphasizes the importance of accounting for acclimatization ability, thermal sensitivity, tolerance and limitations imposed by short- and long-term temperature variability in climate change studies. In a scenario of warming climate, thermal heterogeneity at the microscale will probably create “losers” and “winners” among individuals of a given species, and will likely shape abundance patterns of local populations and thus geographic ranges of species. This study demonstrated that sun exposed limpets frequently face temperatures which can be higher than the *ABT* and very close to the *HCT* during summer low tides.

Living under prolonged sublethal thermal conditions during emersion is energetically costly and can impact the bioenergetic budget and thus the performance (growth, reproduction, foraging ability, Pörtner et al., 2006) and fitness (Somero, 2002, 2010) of these organisms. This indicates that *P. vulgata* may narrow its high shore distribution to shaded microhabitats or surfaces characterized by low exposure to solar radiations in the future climate (see also Lima et al., 2015). Further studies are required to investigate the potential buffering effect of thermoregulatory behaviours (e.g. shell raising Williams et al., 2005). These behavioral strategies could potentially alleviate the bioenergetics cost of living in a thermally stressful environment by reducing the exposure to temperatures capable of inducing cardiac failure and entry into heat coma (Marshall et al., 2013). *P. vulgata* has previously been shown to keep faithful to its home scar (Lima et al., 2015) during the hottest periods of the day, and therefore individuals do not relocate to a less stressful microhabitat in case of heat stress. Recent field investigation has instead shown that *P. vulgata* can raise its shell to significantly lower its body temperatures in comparison to surrounding substrate temperature during heat stress maintaining body temperature at a maximum of 40 °C (maximal temperature inducing cardiac failure in the present study) whilst substrate temperature could reach 42 °C (Chapperon, unpub. data). Therefore, studies integrating thermal dependent physiology, behavior, and biochemical responses at the molecular level are highly needed for a mechanistic understanding of heat stress responses in ectotherms and to predict their vulnerability and distribution patterns in the warming climate.

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Appendix A. Supplementary information

Supplementary data associated with this article can be found in the online version at <http://dx.doi.org/10.1016/j.jtherbio.2016.03.002>.

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