INTRODUCTION

Many biological studies focus on integrating physiological mechanisms such as metabolic performances and thermal tolerances within a broader ecological context. This is typically undertaken to improve the accuracy of predictions regarding the impact of environmental changes on the organism in question (Calosi et al. 2016). However, such research is too often based on broad-scale trends, thereby flattening both microhabitat complexity and the capacity of the organisms to buffer their responses (Hel- muth et al. 2014).

Indeed, different species may show highly variable response times (Kingsolver & Woods 2016). Therefore, understanding the variety of strategies adopted by a species might allow for a better comprehension of the complex mechanisms involved in determining their sensitivity to climatic change (Magozzi & Calosi 2015) and their response to climate-related stressors (Pörtner 2010, Helmuth et al. 2014).

These aspects are particularly crucial when considering benthic sessile species, which are generally more vulnerable to the effects of climate change. This is largely due to their reduced motility, which does not allow them to alter their position in response to changing environmental conditions. To date, relatively little is known about which adaptation strategies organisms select to store energy and why particular responses to stressors remain similar in taxa which occur in habitats across a range of temperatures (Knight 2011, Marshall et al. 2011).

To address this issue, the present study explores the influence of 2 important stressors in shaping the ecological performance of tunicates, using the solitary ascidian Styela plicata (Lesueur, 1823) as a model species.
cies. *S. plicata* is a eurythermal species, widely distributed in tropical and warm-temperate marine and brackish waters (Thiyagarajan & Qian 2003). The species is believed to have originated from the northwestern Pacific. It is a suspension feeder, capturing particles from the water which are then transported on a mucus net present across the branchial basket (Riisgard & Larsen 2010). *S. plicata* benefits from the complex folding of its gill in that its physical size is reduced and it has a greater capacity for water filtration than other ascidians (e.g. Draughon 2010). The minimum size limitation for particulate removal by *S. plicata* is sufficiently small to trap both bacterial and phytoplankton communities (range 0.6 to 20 µm; e.g. Flood & Fiala-Medioni 1981). Although maximum size limits have not been established, glass beads with diameter ranging between 0.5 and 2 mm can be retained by *S. plicata*, suggesting that it may also affect the ichthyofaunal community by removing sperm, eggs or embryos (Draughon 2010).

The Ascidiacea are important because of their unique evolutionary position. Indeed, genomic evidence has recently placed tunicates as the closest link between chordates and ancestral non-chordate deuterostomes (Delsuc et al. 2006). Moreover, their morphology and functional diversity make ascidians good candidates for invading new areas. As such, this includes the potential for negative impacts in the form of physical, ecological, and economic damage (Aldred & Clare 2014). Regardless of the studied species, however, understanding how critical biological processes change under fluctuating environmental conditions is crucial. Responses can include alterations in the consumer’s food intake, assimilation, and metabolic maintenance. Such information can be used to predict how much energy is assimilated and assigned to different needs (growth, development, and reproduction) so that the organisms can resist or adapt to environmental changes in particular habitats (Helmuth et al. 2014).

The aim of the present study was to investigate the ability of *S. plicata* to cope with variations in food availability and environmental temperatures. These parameters were examined separately to test the species’ adaptive capability to colonise different habitats and to perform ecosystem functions such as maintenance of water quality.

**MATERIALS AND METHODS**

Throughout April and May 2015, about 500 individuals of the sea squirt *Styela plicata* were collected from Cala (38° 12' 42” N, 13° 37’ 20” E), a harbour area located within the Gulf of Palermo (Sicily, Italy). This species colonises most of the mooring lines at the study site located at depths of 1 to 3 m. Animals ranging between 3 and 7 cm in length and between 7 and 40 g in wet mass were selected and transferred to the laboratory within 1 h, where they were immediately cleaned of any epibionts by means of a scalpel and placed in a container filled with local seawater for at least 2 h. To minimise any physiological disturbance caused by collection and by floating at the surface of the water, individuals were fixed with acrylic glue to the external side of a 65 mm plastic Petri dish, previously drilled. No mortality occurred during this procedure. Subsequently, animals were maintained for 2 d in 60 l aquaria in aerated and filtered (0.45 µm) seawater at the temperature recorded during sampling (hourly water temperature from 15 to 17°C; www.mareografico.it). During this acclimation period, salinity was monitored and tunicates fed with the microalgae *Isochrysis galbana* twice daily. Prior to starting the physiological measurements, the animals were starved for at least 24 h. The above laboratory procedures were designed to evaluate the physiological performance of individual *S. plicata* in terms of respiration rate (RR, µmol h⁻¹) at varying temperatures and feeding behaviour.

RR was measured for each individual for 1 h. Eight different replicates were used for each temperature level. Single individuals were placed in glass respirometric chambers (500 ml) containing filtered air-saturated seawater. Temperatures were kept stable by means of a thermal bath and monitored throughout the recording period. Magnetic stirrer bars ensured water mixing within the chambers, while oxygen reduction was measured by means of 2 optical oxygen meters (Pyro Science Firesting O2). The latter enabled us to simultaneously monitor 8 different chambers for each experimental temperature through the use of specific software (Pyro Science). To evaluate the thermal tolerance of *S. plicata*, RR was measured at 17 different temperature levels (6, 8, 10, 12, 15, 18, 21, 24, 27, 29, 31, 33, 35, 37, 39, 40, and 41°C). Starting from the water acclimation temperature, the temperature in each group was increased/decreased to the next level at a rate of 1°C per hour (e.g. Fusi et al. 2015).

Clearance rates (CR, l h⁻¹), measured as the volume of water cleared of algal cells per hour, were determined using a static system (Widdows & Staff 2006). The decrease in algal cell density in each experimental beaker was monitored after a given time interval. Feeding and assimilation rates were determined for...
8 different food concentrations (0.3, 0.6, 1.4, 2.2, 3.1, 4.4, 8.6, and 15.8 µg l\(^{-1}\) chl \(a\)), using 10 different animals in each treatment.

Specifically, after about 30 min to allow the organism to recover from manipulation, aliquots of the algal culture (\(I.\) galbana) were added to each beaker. Then, 20 ml subsamples were collected every 30 min for 2 h and the cell density measured using a Z2 Coulter Counter (Beckman Coulter) fitted with a 100 µm orifice tube. CR was calculated as Vol × (\(C_{1} – C_{2}\) / time, where Vol is the volume of water, and \(C_{1}\) and \(C_{2}\) are the mean cell concentrations at the beginning and end of each time increment (hours).

At the end of the CR measurements, specimens of \(S.\) plicata were singularly placed (i.e. 1 ind. beaker\(^{-1}\)) into new 1 l beakers containing filtered seawater and left undisturbed overnight to produce faeces. Accumulated faeces and food provided for each treatment replicate were collected by filtering the water from each beaker with GF/C filters, combusted and preweighed. The efficiency of food assimilation (AE) was calculated using the Conover (1966) ratio, \(AE = (F - E) / [(1 - E)F]\), where \(F\) and \(E\) are the ash-free dry weight (DW):DW ratio of the food ingested and faeces, respectively. The corresponding weight of the organic material combusted was obtained by drying the faeces and algal mass at 105°C for 24 h. This was then weighed and ashed at 450°C for 4 h before being weighed a second time. Finally, 3 other replicates of every food treatment were filtered onto GF/F filters and used to quantify the concentration of chlorophyll \(a\) (chl \(a\)), which served as a descriptor of species feeding. Spectrophotometric analyses of chl \(a\) were carried out according to Lorenzen & Jeffrey (1980) by extracting pigments using 90% acetone (24 h in the dark at 4°C).

**Statistical analysis**

Differences in metabolic rates and feeding responses under the varying experimental conditions were tested separately (i.e. without any combinations of both factors) and were analysed by means of ANOVA (Underwood 1997). Temperature (17 levels) and food concentration (8 levels) were considered fixed factors, using 8 specimens for temperature levels and 10 specimens for food concentrations, and treated as replicates. The assumption of homogeneity of variance was tested \(a\) priori by means of the Cochran’s test. The Student-Newman-Keuls (SNK) test allowed the appropriate means comparison (Underwood 1997). When no homogeneous variances were rendered for any type of transformation, as in the case of AE, the significance level was set at 0.01 instead of 0.05. This reduced the possibility of a Type I error (Ruiz et al. 2010, Sarà et al. 2013). ANOVA was carried out using the GMAV software (version 5.0).

**RESULTS AND DISCUSSION**

Across the temperature ranges between 6°C and 41°C, the amount of oxygen consumed mainly varied between \((\text{mean} \pm \text{SE})\) 1.25 ± 0.27 µmol h\(^{-1}\) g\(^{-1}\) DW (estimated at 6°C) and 16.49 ± 1.85 µmol h\(^{-1}\) g\(^{-1}\) DW (at 29°C). The only exception was recorded at 40 and 41°C, where the average metabolic rate was 2 times higher \((32.99 \pm 5.17 \mu \text{mol h}^{-1} \text{g}^{-1} \text{DW})\) (Fig. 1). ANOVA revealed a significant effect of temperature on the metabolic rate of \(S.\) plicata \((p < 0.001;\) Table S1 in the Supplement at www.int-res.com/articles/supp/b026p179_supp.pdf), with significant differences recorded among the rates estimated at lower (6°C and 8°C) and higher (40°C and 41°C) temperatures (Table S2).

\(S.\) plicata is commonly found in harbours, salt marsh and marine habitats in both warm and temperate waters (the Pacific, Indian and Atlantic oceans, and the Mediterranean Sea). Thus, the width of the thermal tolerance window observed in the present study overlaps with this distributional pattern. \(S.\) pli-
cata also exhibits the ability to maintain relatively low metabolic rates within a broad range of environmental temperatures. Again, the tested specimens showed an active metabolism under almost all temperatures, with thermal limits well within the range reported in the literature. The lowest temperature of metabolic functioning was 6°C (Fig. 1), a temperature previously reported as lethal for *S. plicata* populations within the Venice Lagoon (Sabbadin 1957). However this figure is slightly lower than the 10°C more recently reported as the lower boundary of its thermal range (Thiyagarajan & Qian 2003). In our study, pairwise comparison showed that *S. plicata* remained the same at temperatures between 21°C and 39°C, suggesting that metabolic maintenance made no excessive demands on the energetic trade-off of the species. This partially confirms the observations of Fisher (1977), who suggested that a lack of larval settlement may not be a consequence of energetic deficiency due to elevated summer temperatures, but rather reflect the contribution of other different local factors. The ability to depress metabolic performance, resulting in a thermally insensitive response (sensu Verberk et al. 2016), could instead be viewed as an adaptive strategy. Thus, individuals maintain an aerobic scope to contextually cope with minor damages over a broad range of temperatures, for example, by increasing heat shock protein (HSP) activity. However prolonged exposure to stress can lead to metabolic failure in a relatively short time. In the higher temperature treatments, *S. plicata* individuals consumed almost all available oxygen in less than 1 h down to concentration of 3 µmol l⁻¹. At this point, the aerobic scope for performance increase was progressively reduced, as represented by the plateau in the recorded metabolic rates. This response is related to the transition from the optimal to the pejus (sensu Sokolova et al. 2012) temperature range in aquatic invertebrates (Giomi & Pörtner 2013), reflecting cascades of molecular responses activated by heat stress, and higher energetic costs. At lower temperatures tested, water oxygen concentrations decreased on average between 3.2 and 55.5% recorded at 6 and 29°C, respectively.

Our results also confirm the considerable ability to assimilate food efficiently when resources are limited. Significant differences in mean CRs were found between individuals fed with different algal concentrations (Tables S3 & S4, Fig. 2a). CR values lower than 1 l h⁻¹ g⁻¹ DW were recorded at algal concentrations between (mean ± SE) 0.3 ± 0.03 and 2.24 ± 0.03 µg l⁻¹ chl a, but increased to 1.35 ± 0.10 l h⁻¹ g⁻¹ DW at concentrations higher than 3 µg l⁻¹.

Maximum CRs (mean ± SE) of 1.53 ± 0.17 l h⁻¹ g⁻¹ DW and 1.50 ± 0.26 l h⁻¹ g⁻¹ DW were calculated at food concentrations of 3.1 and 8.6 µg l⁻¹, respectively. These figures are comparable with those of Draughon (2010), who reported an average CR of 1.66 l h⁻¹ g⁻¹ DW for *S. plicata* fed with *Nannochloropsis*, a micro-algae similar in cell size (2 to 4 µm) to that used here (*Isochrysis galbana*, 2 to 5 µm). The variation in CR estimates between the studies may reflect the difference in the size of the organisms used to carry out the feeding experiments. Those employed in the present study were smaller than those collected by Draughon (2010), where wet weights ranged between 20 and 50 g. The tendency of CR to increase even when resources are scarce (e.g. oligotrophic waters with <1 µg l⁻¹ chl a) confirms that ascidians have adapted to environments with low concentrations of particles.
However, a constant ingestion rate was recorded over a certain threshold, probably as a result of gut satiation (Petersen et al. 1999). In other words, the ingestion rate of *S. plicata* at most food densities is described by the Holling (1959) Type II functional response. Over the threshold of 8.6 µg l–1, specimens decreased their filtration rate by increasing saturation in a strictly monotonic manner (Jeschke et al. 2004). The food threshold detected in the present study is also in line with concentrations reported in previous studies (8 to12 µg l–1 chl a; Petersen & Riisgard 1992).

In contrast, food concentrations did not affect the efficiency with which *S. plicata* assimilated the organic material; no significant differences were detected between treatments (Table S3). The average AE value was (mean ± SE) 76 ± 10% (Fig. 2b), ranging between 0.68 ± 0.03% and 0.82 ± 0.01% at intermediate and low food densities, respectively. These results confirm the great ability of this group to retain microalgal cells (particle sizes ranging between 1 and 7 µm; Randlov & Riisgard 1979) and to not differentiate between the removal of inert, inorganic, and organic particles (Draughon 2010). Such efficient feeding strategy management also suggests that this species has potential to be used as a biological filter in an aquatic environment. In addition to reducing bacterial and microalgal blooms in estuarine waters (e.g. Draughon 2010), ascidian species such as *S. plicata* and *Ciona intestinalis* can also act as hosts to several different types of organisms, and serve as sentinel organisms in the bioremediation of heavy metals (Abdul Jaffar Ali et al. 2015).

In conclusion, *S. plicata* are able to efficiently manage their own metabolic trade-off. This species’ ecophysiological performance may thus underpin its successful colonization of new areas with different environmental characteristics (Bates et al. 2013). In addition, it may also provide evidence of the evolution of temperature-insensitive metabolisms to save energy across part of the thermal window (Marshall et al. 2013). Maintaining elevated retention efficiency and constancy of metabolic costs within a wide range of temperatures may reduce the risk of mortality when an organism’s metabolic demand (which is temperature-dependent) cannot be met by supply. Furthermore, this may also contextually confer a competitive advantage for this introduced (sensu Clarke-Murray et al. 2012) species, which is capable of causing extreme modification of the structure of coastal habitats (e.g. Sutherland 1978). Thus, *S. plicata*‘s life history traits (e.g. fast growth and hermaphroditic reproduction), lower resting maintenance costs and ecophysiological features such as a wide range in diet, likely explain its spread and establishment into new areas. Understanding the mechanisms of thermal adaptation and response to stressors in *S. plicata* will help improve predictions of physiological and evolutionary mechanisms driving current and future distributions of this species (Woodin et al. 2013).

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