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Compared stress tolerance to short-term exposure in native and invasive tunicates from the NE Atlantic: when the invader performs better

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Abstract

The combined impact of invasive species and climate change threatens natural systems worldwide, often facilitating the expansion of harmful invasive species. It is imperative to understand the mechanisms behind why species become invasive and widespread. Traditionally, it is thought that invasive species have greater tolerances to a wider array of environmental conditions than natives. We, therefore, tested the hypothesis that invasive species are more tolerant to the effects of short-term exposure to temperature and salinity stress. Using unifactorial experiments, we compared the tolerances of two common fouling NE Atlantic ascidians, the native *Ciona intestinalis* and the invasive *Styela clava*, to increased temperature and decreased salinity. We measured lethal and behavioural responses affecting 50% of populations to give an indication of the tolerance limits for temperature (LT₅₀) and salinity (EC₅₀), and respiration rate to give an indication of the change in metabolic response. The invasive *S. clava* was more tolerant to increased stress (LT₅₀=29.5 °C, EC₅₀=19.5) compared with *C. intestinalis* (LT₅₀=27.0 °C, EC₅₀=22.7), whereas both species displayed similar metabolic responses observed through increased respiration rates. This study is among the first to experimentally determine limits for temperature and hyposalinity stress for either species and supports the hypothesis that the invader performs better under extreme conditions. Future environmental changes caused by events such as heat waves and climate change could push species to the edge of their physiological limits, potentially facilitating competitive shifts between native and invasive species.

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Introduction

On a global scale, the impact of invasive species is paramount, causing detrimental ecological impacts to ecosystems and their native communities (Mack et al. 2000; Butchart et al. 2011). In combination with climate change, the impacts of invasive species threaten biodiversity worldwide, potentially causing wide-ranging effects. These include impacting native species abundances and distributions, local extinctions, alteration of vital ecosystem functions and services, and significant economic impacts (Pimentel et al. 2005; Halpern et al. 2008; Hellmann et al. 2008; Pejchar and Mooney 2009; Butchart et al. 2011). Furthermore, these two drivers of change often interact with climate change enhancing the spread of non-native species (Dukes and Mooney 1999; Hellmann et al. 2008).

Evidence supports the hypothesis that shipping and hull fouling is a major vector in transporting marine species worldwide (Clarke Murray et al. 2012; Peters et al. 2017). What is less clear are the mechanisms employed by invasive species to enable them to become so pervasive. Generally, for invasive species perceived higher physiological tolerances to a range environmental conditions (e.g. temperature, salinity, pollution levels) is thought to enable greater competitive ability over native species, helping to facilitate successful settlement, establishment of populations, and further spread (Lenz et al. 2011; Zerebecki and Sorte 2011; Lejeusne et al. 2014; Lagos et al. 2017).

Invasive species often cause serious detrimental impacts. Of the biofouling taxa, tunicates are a major concern. They can cause smothering of aquaculture facilities and species, damage to structures, increased drag on propellers resulting in reduced efficiencies, and significant cleaning costs (Aldred and Clare 2014). For example, the highly invasive tunicate Styela clava presents a significant global risk. Its introduction in the Gulf of St Lawrence, Canada, resulted in a 50% loss to shellfish aquaculture industries (Colautti et al. 2006). In the present study, we, therefore, address the tolerance of S. clava in comparison to the NE Atlantic native tunicate Ciona intestinalis, which is also considered invasive in other parts of the world and causes significant economic impacts (e.g. Colautti et al. 2006; Therriault and Herborg 2008a). It is imperative that we understand the physiological mechanism behind these species invasion success.

Recently, C. intestinalis has undergone taxonomic reevaluation. Formerly, at least two cryptic species, Types A and B, made up this species complex and are now known as C. robusta and C. intestinalis, respectively (Brunetti et al. 2015). Ciona robusta prefers warmer waters, is native to northwest Pacific and is introduced worldwide including Europe. Conversely, C. intestinalis is considered a cold water species native to northwestern Europe, from northern Portugal to Norway (Bouchemousse et al. 2016a). This species is also considered invasive in China (Zhan et al. 2010) and the northeastern American coastline (Therriault and Herborg 2008b). However, recent genetic evidence suggests an amphi-Atlantic native distribution (Bouchemousse et al. 2016a). Prior to taxonomic re-evaluation, this species was considered to tolerate temperatures exceeding 30 °C (Dybern 1965; Marin et al. 1987; Carver et al. 2006); however, the highest temperature tolerances were attributed to specimens found in the Mediterranean where C. robusta is pervasive (Zhan et al. 2010). So virtually no information is available on the tolerance of C. intestinalis to thermal stress. Similarly, previous data regarding the salinity tolerance of this species should be carefully interpreted considering taxonomic re-evaluation. Generally C. intestinalis is considered a euryhaline species and, among populations within this species native range, it has been observed to tolerate salinities under 12 (Dybern 1967; Carver et al. 2006).

By contrast *S. clava* is native to the northwest Pacific but has spread worldwide including to Europe, North America and Australasia (Lützen 1999; Davis and Davis 2008). Within these locations, its settlement is known to result in significant economic impacts to aquaculture; in Canada alone this was estimated between CA\$ 34 and 88 millions per year (Colautti et al. 2006). Within Europe, it was first recorded in 1953 (designated as Styela mammiculata) in the Lynher Estuary, Plymouth, UK (Carlisle 1954). It then started expanding across northwestern Europe during the following decades (Lützen 1999) and was first recorded in the Mediterranean by the mid-2000s (Davis and Davis 2008). Major vectors of this spread can be attributed to human-mediated transport on ship hulls and within ballast water whereas natural dispersion is responsible for spread to neighbouring sites only (Davis et al. 2007). Experimental and observational studies of specimens located outside of this species native range show that S. clava settles in locations experiencing temperatures ranging from - 2 °C to in excess of 23 °C (Buizer 1980; Davis and Davis 2008). However, there is limited evidence to suggest its maximum tolerance. In addition, these studies have also indicated a preference for warm waters, showing that for successful reproduction and settlement of this species ambient water temperatures must exceed 16 °C for several days (Davis et al. 2007). This species also has limited osmoregulation capabilities, preferring salinities above 20 (Davis et al. 2007; Davis and Davis 2008). However, it has been known to survive lower salinities by closing siphons for prolonged periods (Sims 1984; Lützen 1999).

Understanding the tolerance of species to environmental conditions is the key to understanding their potential spread. Regarding the two species of interest here, while they are common in urbanised habitats, often dominating fouling communities (Lambert and Lambert 1998; Gittenberger and Van Der Stelt 2011), their tolerance to temperatures and salinities is not widely understood, particularly with regard to the metabolic impacts of these stressors. Furthermore, the recent taxonomic re-evaluation of C. intestinalis calls to question previous evidence of tolerance for this species. We, therefore, conducted a series of short-term (24 h) experiments examining the impact of increased temperatures and decreased salinities on S. clava and C. intestinalis, evaluating survival and metabolic response in terms of respiration. These were used to determine upper temperature and lower salinity tolerance limits. These limits are known to be strongly correlated with the range of conditions a species would be expected to be able to tolerate under natural conditions (Zerebecki and Sorte 2011; Kelley 2014); therefore, these upper limits give an indication of the capacity of these species to be able to survive extreme conditions. We hypothesised that the invasive S. clava would display a greater tolerance to short-term hyposalinity and increased temperature in comparison to the native C. intestinalis, both in terms of survival and metabolic response.

Methodology

Collection and maintenance of organisms

Ciona intestinalis and S. clava were collected from the Château Marina in Brest, France (48°22'44"N, 4°29'21"W) in June 2017. Seawater temperature was continually monitored at three locations within the marina using HOBO (UA-002-64) data loggers, deployed between the 26 March and 4 October 2017 and attached to floating structures (Supplementary Material). Individuals were found in high abundances attached to artificial substrates (pontoons, ropes, metal pillars). Species were collected from under pontoons by SCUBA diving, carefully scraping and removing adults from structures located within the centre of the marina. Organisms were subsequently transported to the Roscoff Biological Station in seawater where they were cleaned of epiphytes and epibionts and placed overnight in running ambient temperature seawater aquariums. Specimens were then transferred into controlled temperature tanks with renewed seawater inflow for long-term storage. Based upon water conditions from which they were taken and temperatures appropriate to maintain live animals, specimens were kept at 16 °C. Every 2 days, specimens were fed a 1:2 (by cell count) mixture of the algae Chaetoceros gracilis (IFREMER strain from Argenton) and Isochrysis galbana (Tahitian strain from Roscoff Culture Collection) totalling approximately 4×10^8 cells L⁻¹. All specimens were acclimated to these conditions for at least 72 h prior to experimentation.

Experimental setup

The same experimental procedure was used for both the temperature and salinity experiments. During each experiment, experimental units (EUs) consisted of 2 individuals of a single species placed in a 1.8-L tank supplied with constant aeration. In total each experiment used 40 EUs—2 species $\times 5$ treatments (levels of salinity or temperature) $\times 4$ replicates for each treatment–species combination. This equated to four EUs per treatment per species which were spread among five large temperature-controlled aquaria (Fig. 1); the heaters used maintained temperature ± 0.5 °C from the target.

After 24 h, each EU was assessed for mortality based upon the number of live and dead individuals within the tank (n=4 EUs per treatment per species). In addition, one individual was removed from each EU to measure respiration. Hence, four individuals were used per treatment per species for respiration (n=4). Specimens used in the respiration measurements were selected based upon their visual size: approximately 5–8 cm for *C. intestinalis* and 8–11 cm for *S. clava*. To determine mortality, three criteria were assessed: contraction (siphon or body), colour change, and loss of response to gentle touching of the siphons.

Temperature experiment

Temperature treatments were as follows: 16 (control: stock tank temperature), 20, 24, 28 and 32 °C. Desired temperatures were achieved by raising the water temperature within the five large temperature-controlled aquaria (Fig. 1). After placing individuals within tanks and starting from 16 °C, temperatures were raised by 1 °C every 15 min. Once the desired temperature was reached, tanks were left for 24 h.



Fig. 1 Experimental setup of the temperature experiment. Each experimental unit (EU; small square) contains two individual specimens of the same species (circles: *Ciona intestinalis* light grey, *Styela clava* dark grey). EUs are placed inside larger temperature-controlled

tanks (large rectangle) filled with freshwater to homogenise and maintain the temperature within each EU over the 24-h experimental duration

To assess respiration, acrylic incubation chambers (600 mL) were utilised using methods from Noisette et al. (2016). Each chamber was filled with seawater that was equilibrated to the same temperature used in the corresponding treatment. One individual from each EU was taken and placed in a single chamber (n = 4 per treatment per species). A magnetic stirrer was placed within each chamber and the chambers were positioned on a waterproof stirring plate. The plate was kept underwater in an additional temperaturecontrolled aquarium, equilibrated to the corresponding treatment temperature. The stirring plate allowed six chambers to be incubated simultaneously. In addition, six control chambers were used for each temperature to correct for fluxes due to microbial activity within the seawater in each of the treatments. Using a fibre-optic system and reactive oxygen spots on the chambers (FIBOX 3, PreSens, Regensburg, Germany), oxygen was measured at the start and end of an incubation period which lasted for 30 min. Respiration (R;in μ mol O₂ g⁻¹ DW h⁻¹) was calculated using the following equation:

$$R = \frac{\Delta O_2 \times V}{\Delta t \times DW},$$

where ΔO_2 (in µmol $O_2 L^{-1}$) is the difference in oxygen concentration between the start and end of the incubation, V is the measured volume of each chamber minus the volume of the specimen, Δt is the incubation time (hours), and DW is the dry weight (g) of the specimen and obtained by placing specimens in a drying oven for 48 h at 60 °C.

Salinity experiment

To obtain the desired salinities, distilled water was added to seawater and checked using a Fisher Scientific[™] Traceable[™] salinometer. Salinity treatments were as follows: 35 (control: natural seawater), 28, 24, 21 and 17. The lower values were chosen based upon the ranges of salinity tolerance observed in other studies (Shumway 1978; Carver et al. 2006; Clarke and Therriault 2007; Davis et al. 2007) and on pilot studies where mortalities occurred between salinities of 14–21. Each EU contained two individuals which were placed directly from natural seawater into altered salinity water. EUs were randomly dispersed amongst the five large temperature-controlled aquaria to prevent fluctuations in temperature over the experimental duration.

This experiment was independent of the temperature experiment and used different apparatus for calculating oxygen concentration. The experimental setup follows the methodology used in the temperature experiment, the differences being the chambers used for incubation and the device used to measure oxygen within the water. The equation for calculating respiration remains the same. In this experiment, incubations were conducted within hermetically sealed glass chambers of two sizes (200 and 500 mL). Due to the smaller *C. intestinalis* used in this experiment, a smaller size of incubation chamber (200 mL) was necessary to allow an appreciable change in oxygen to be able to calculate respiration accurately. This chamber was too small for *S. clava* which was, therefore, incubated in the larger (500 mL) chamber. Oxygen was measured using an oxygen probe (Hach-Lange LDO101) prior to sealing the chambers (incubation start) and following the incubation (approximately 30 min), at which point the seal was broken and the oxygen measured immediately.

Data analysis

Mortality data were used to calculate the temperature that was lethal to 50% of the population (LT_{50}). For the salinity experiments, at the range used, a behavioural response was observed rather than a mortal response; the effective concentration (EC_{50}) was, therefore, calculated and defined as the salinity at which 50% of individuals displayed a response. Binomial regression models using the probit link function were used to model the response and to produce mortality curves. Using these models, the LT_{50} and EC_{50} and corresponding confidence intervals were calculated. All analyses were completed using the statistical programme R, version 3.22 (R Core Team 2015) and visualised using the ggplot2 package (Wickham 2009).

Two-factor ANOVA was used to analyse the respiration data, which, depending on the experiment, used either the temperature or salinity and species as fixed factors. To facilitate a balanced statistical design in the temperature experiment, the highest temperatures (28 and 32 °C) were not included in the analysis due to high mortality. To explore interaction terms and explore the impact of temperature on individual species (thus incorporating excluded data for higher temperatures where data were available), single-factor ANOVA and Tukey's HSD *a posteriori* comparisons (where appropriate) were performed for each species. Residuals of the model were checked for normality and homogeneity of variances; the salinity data set was log transformed to meet normality and homogeneity of variances criteria.

Results

Temperature experiment

After 24 h, increased temperatures had a significant effect on the mortality of both *C. intestinalis* and *S. clava* (Fig. 2a); in both species 100% mortality occurred within the highest temperature treatment (32 °C). There was a small difference in the modelled LT_{50} values observed in both species. The invasive *S. clava* displayed higher tolerance to increased

Fig. 2 a Mean probability of mortality $(\pm SE)$ and **b** respiration (μ mol O₂ g⁻¹ dry weight h⁻¹) of Ciona intestinalis and Styela clava exposed to increased temperatures for 24 h (n=4). Mortality curves (a) were produced using Probit analysis and indicate the probability of a species mortality at a given temperature; 50% mortality is indicated for each species (LT₅₀). For respiration (**b**), bars represent mean respiration $(\pm SE)$ and circles are individual data points, as within the ANOVA there were significant main effects, letters indicate significant differences within the temperature factor only, as determined by a posteriori analyses (28 °C was not included in the analysis due to lack of replication)



temperatures ($LT_{50} = 29.5$ °C; CI = 27.1–31.9) than the native *C. intestinalis* ($LT_{50} = 27.0$ °C; CI = 25.5–28.5).

In the analysis of respiration (Table 1), there was no significant interaction term between species and temperature treatments indicating respiration in both species was affected similarly by increased temperatures. Overall respiration was significantly lower in *S. clava* compared to *C. intestinalis*. Temperatures above 16 °C resulted in the highest respiration for both species (Fig. 2b). There was 100% mortality at 32 °C, and only 1 surviving *C. intestinalis* at 28 °C on which respiration could be measured; therefore, only the first 3 temperature treatments were considered in the twofactor ANOVA (Table 1). Increased temperatures resulted in significantly higher respiration in comparison to the 16 °C temperature treatment. There were no differences between the measured respiration at 20 °C and 24 °C for *C. intestinalis* or at 28 °C in *S. clava* (Fig. 2b).

Salinity experiment

There was a minimal effect of lowered salinities on mortality at the ranges tested. Only two specimens were clearly identified as dead, meeting all three criteria of mortality as defined in the methodology. There was, however, a behavioural response whereby specimens no longer responded to touching their siphons. In addition to being unresponsive,

 Table 1
 Two-factor
 ANOVA examining the effects of temperature

 (A) and salinity (B) on two species of ascidian—the native *Ciona intestinalis* and the invasive *Styela clava*—using species and the stressor (either temperature or salinity) as fixed factors

Factor	df	MS	F	р
(A) Temperature				
Temperature	2	636.1	27.13	< 0.001
Species	1	126.9	5.41	0.031
Temperature \times species	2	10.1	0.43	0.656
Residuals	19	23.4		
(B) Salinity				
Salinity	4	0.6	4.07	0.009
Species	1	3.99	27.17	< 0.001
Salinity \times species	4	0.15	1.01	0.416
Residuals	30	0.15		

Effects of temperature were examined at three levels (16, 20 and 24 °C) and salinity at five levels (17, 21, 24, 28 and 35). Bold type indicates a significant effect at p < 0.05; n = 4 for all treatments

the siphons of *S. clava* specimens were continually open (not contracted), whereas in *C. intestinalis*, specimens were extremely contracted. Therefore, in the following analysis, this behavioural response has been modelled, calculating the effective salinity impacting 50% of the population (EC_{50}).

Both species displayed a tolerance to decreased salinities; however, after 24 h, all individuals in the lowest salinity treatment displayed the described behavioural response (Fig. 3a). *Styela clava* displayed the greatest tolerance to decreased salinity with an EC₅₀ of 19.5 (CI = 17.3–21.6) whereas *C. intestinalis* displayed an EC₅₀ of 22.7 (CI = 21.6–23.8).

Within the two-factor model analysing the impacts of salinity on respiration (Table 1), there was no significant interaction term, while the main effects were significant. Within the salinity experiment there was a large difference in respiration rates observed between the two species. Similar to the temperature experiment, overall higher respiration by body mass was observed in *C. intestinalis*. Salinity had a significant effect on both species (Table 1) whereby respiration rate was lower in natural seawater (salinity 35) in comparison to salinities less than 28 (Fig. 3b).

Discussion

In experiments examining the tolerance of two common fouling species to increased temperatures and decreased salinities, the invasive *S. clava* and the native *C. intestinalis* both displayed high tolerances to short-term (24 h) exposure. Comparatively, while in both species the respiration rates responded similar to the altered salinities or temperatures, *S. clava* displayed tolerances greater than that of *C. intestinalis* in terms of displaying mortal (LT₅₀) and behavioural responses (EC₅₀). This is in agreement with other studies stating invasive species have a wider tolerance range which, among other responses, impacts survival and respiration (Lenz et al. 2011; Zerebecki and Sorte 2011; Lejeusne et al. 2014). Worldwide, both species are commonly found fouling urbanised habitats such as marinas and are considered invasive species (Davis and Davis 2008; Zhan et al. 2010). As such, human-induced dispersal and climate change influencing temperature and salinities have the potential to influence worldwide distribution patterns and range expansions of these species (De Rivera et al. 2011; Rius et al. 2014). This study offers empirical evidence of the versatility of these ascidians, particularly regarding S. clava which is a highly prolific invader. This species was shown to tolerate a wide range of environmental conditions, therefore, facilitating its spread and the threat of future invasion success.

Thermal tolerance

In the mortality analysis, the invasive S. clava was able to tolerate and survive at higher temperatures than the European Atlantic native C. intestinalis ($LT_{50} = 29.5$ °C and 27 °C, respectively). In terms of the respiratory response, both species were affected similarly by increased temperature. This complements previous experimental studies documenting that invasive species temperature tolerances are greater than those of comparative native species (Zerebecki and Sorte 2011; Kelley 2014; Lejeusne et al. 2014). Furthermore, we empirically show upper temperature limits to acute thermal stress (LT₅₀), providing information on each species thermal niche. Higher upper thermal tolerance limits are known to be a strong indicator of the eurythermality of a species (Kelley 2014). When assessing multiple species, these upper limits can, therefore, give a good indication of comparative tolerances of species, thus relating to how species would cope with environmental changes in the wild. Based on geographic ranges, C. intestinalis is documented to be found in temperatures ranging from subzero to up to 24 °C (Dybern 1965; Carver et al. 2006; Vercaemer et al. 2011). However, the recent taxonomic separation of C. intestinalis (previously C. intestinalis type B) from its congener C. robusta (type A) calls to question tolerance experiments conducted prior to its separation. By comparison, previous monitoring has shown that S. clava typically establishes invasive populations in locations where temperatures range from subzero to in excess of 23 °C (Buizer 1980; Davis and Davis 2008); however, there is less information on a maximum temperature for this species. In an example of known populations where sea surface temperatures have exceeded these limits, Davis and Davis (2010) noted that in Sète, French Mediterranean, extreme summer temperatures reached 29.1 °C in 2006. While this is only slightly lower

Fig. 3 a Mean probability of the loss of response $(\pm SE)$ and ${\boldsymbol b}$ respiration (µmol $O_2 \ g^{-1}$ dry weight h⁻¹) of Ciona intestinalis and Styela clava exposed to decreased salinities for 24 h (n=4). Response curves (a) were produced using Probit analysis and indicate the probability of a species losing responsiveness to stimuli at a given salinity; 50% of the populations' response loss is indicated for each species (EC₅₀). For respiration (**b**), bars represent mean respiration $(\pm SE)$ and circles are individual data points, as within the ANOVA there were significant main effects, letters indicate significant differences within the salinity factor, as determined by a posteriori analyses



than the LT_{50} we observed here, the survival of this population in Sète was originally theorised to be due to refuge habitats at lower (and cooler: 26.6 °C) depths (Davis and Davis 2010). We show that *S. clava* could survive these extreme events, at least for the short term.

The IPCC predicts that climate change is expected to increase global temperatures worldwide by between 1 and 3.7 °C by the end of the century (IPCC 2013) and increase the severity of summer heat waves by up 2 °C in the coming decades (Meehl 2004; Perkins-Kirkpatrick and Gibson 2017). Extreme summer temperatures have already been shown to significantly impact subtidal communities (Lejeusne et al. 2010; Sorte et al. 2010; Smale et al. 2015).

With increasing severity, the observed effects will become more pronounced. At present, in the Château Marina in Brest, France, we observed maximum summer water temperatures of 21 °C and the summer averages approximately 18 °C (Supplementary Material). Future increases in temperature will likely impact fouling communities and could potentially impact invasive and native species differently based upon their respective tolerances (Sorte et al. 2010; Smale et al. 2015).

With respect to *S. clava* and *C. intestinalis*, it is likely that temperatures up to or exceeding the LT_{50} for both species could be detected within Europe by the end of the twenty-first century. This could have significant impacts on the

range of S. clava, causing it to become an even more ubiquitous invader in European waters and worldwide. Already we are seeing this species moving further north (e.g. Cook et al. 2013) but there is potential for it to spread to new locations around the Iberian Peninsula and the Mediterranean (Davis and Davis 2010; Çinar 2016). While ocean warming could facilitate the spread of S. clava, the native C. intestinalis is a cold water species and its hypothetical range expansion due to climate change would be more limited. At present, C. intestinalis is dominant within Brittany where this species occurs in syntopy with C. robusta. It is known that warmer summers in these locations tend to facilitate increased settlement of the invasive C. robusta (Bouchemousse et al. 2016b) and monitoring suggests C. intestinalis has a much lower thermal maxima and is less tolerant to thermal changes than C. robusta (see Bouchemousse et al. 2016a, b). With respect to climate change, increased temperatures could impact physiological mechanisms and result in higher mortality in C. intestinalis, thus decreasing its competitive ability over C. robusta and resulting in a competitive shift between the two species. Given that the tolerance of native species tends to be lower than taxonomically similar invasive species (Lenz et al. 2011; Lejeusne et al. 2014), other species will likely be impacted similar to increased temperatures and result in significant community shifts.

While the survival data relate to thermal maxima, a functional stress response can be observed for both species at lower temperatures. This directly relates to the tolerance of a species and its potential to become an invader outside of its native range. For both species, the rates of respiration increased when temperatures exceeded 20 °C, where it reached its maximum. At temperatures above 20 °C, there was zero or negligible increases in respiration rate. Similar results were observed by Ai-Li et al. (2008) and Kang et al. (2015) who confirm this relationship is consistent among size classes of S. clava. Similar results have also been observed on other physiological responses in C. intestinalis. For example, in populations from Denmark, Petersen and Riisgard (1992) discovered that filtration rates began to decline above 21 °C, whereas at lower temperatures, a linear relationship between filtration rate and increasing temperature was observed. This suggests thermal stress was occurring above these temperatures and complements the present study where a metabolic response is observed and peaks at 20 °C. At higher temperatures, there were no further significant changes in respiration rate; however, given that there was high mortality in our treatments above 24 °C, it is reasonable to suggest temperature is causing a significant stress response, at least for *C. intestinalis*.

In this study, no further increases in metabolic response over 20 °C indicate a reduction in temperature dependency of the metabolism (i.e. at higher temperatures metabolism is not impacted by increasing temperature). While this could be interpreted as an indication that the animal is stressed, measurement of multiple physiological mechanisms should be regarded to fully consider an organism's ability to survive or grow (e.g. scope for growth model: Warren and Davis 1967; Newell and Branch 1980). In other studies of temperature effects on various physiological mechanisms employed by S. clava, temperatures above 24 °C have resulted in declines of feeding and excretion rates, and temperatures of 28 °C have been shown to cause declining function and scope for growth (Jiang et al. 2008; Kang et al. 2015). However, even within this temperature range, S. clava has been shown to be well adapted, with the capacity to maximise its physiological response according to the temperature to meet energy demands (Jiang et al. 2008). The higher tolerance displayed by S. clava and other invasive species is perhaps one of the most significant features that allow them to spread and establish worldwide.

Hyposalinity tolerance

Our results correspond to the tolerances observed within natural ranges of both species. Ciona intestinalis is a euryhaline species, reportedly within its native habitat tolerating salinities ranging from 12 to 40 (Dybern 1967; Carver et al. 2006; Therriault and Herborg 2008b). By contrast, the invasive S. clava is rarely found below salinities of 20 (Lützen 1999; Davis et al. 2007; Davis and Davis 2008). However, this species has been known to survive short durations when subjected to salinities below 20, purportedly surviving for several days by closing siphons (Sims 1984; Lützen 1999). Similar behavioural responses have been observed in C. intestinalis which, in addition, has been shown to cease respiring during times of lowered salinity (Shumway 1978). This behavioural response of closing siphons is, perhaps, related to avoidance mechanisms similar to those employed by other species, such as bivalves closing shell valves (e.g. Davenport 1977) or burrowing behaviour in polychaetes (e.g. Shumway and Davenport 1977). This mechanism was not employed within our experiments for either species; however, for both species, cessation of response to siphonal stimuli did occur, indicating a severe stress response to decreased salinity. In mortality studies on tunicates, this lack of response is often recorded as an endpoint (Sims 1984; Peck et al. 2009; Zerebecki and Sorte 2011; Jofré Madariaga et al. 2014); however, as seen in this study, the lack of a response does not indicate death or cessation of respiration. Other factors should be taken into consideration to assess mortality or fitness of an individual, especially under salinity stress.

To our knowledge, only two previous studies have researched the effects of lowered salinities on the respiration of the two target species (ISI web of science search, accessed March 2018, using keyword combinations of salinitylrespirationloxygen consumption-no further results were found when the search was expanded to include other ascidians). Of these, both studies present results contradictory to our own. While we identified an increase in respiration due to the effects of lowered salinity, Shumway (1978) and Ai-Li et al. (2008) show decreasing salinities caused reductions in the respiration of C. intestinalis and S. clava, respectively. However, these studies focus on shorter duration exposures than the results we present (6 and 2 h, respectively, compared with our 24-h exposure). Longer term subjection to lowered salinities could facilitate acclimation and a response related to longer term exposure. Kinne (1966) proposed four categories describing the varied physiological responses of organisms when exposed to changes in salinity, stating that metabolic rates will either be "(1) increased by subnormal salinities, and/or reduced in supranormal salinities, (2) increased both in sub- and supra-normal salinities, (3) reduced both in sub- and supra-normal salinities, (4) essentially unaffected". Within this study, our results correspond to categories 1 and 2 describing increases in metabolic response caused by subnormal salinities. This is a response typical of other euryhaline species (e.g. Kinne 1966; Roast et al. 1999; Ern et al. 2014). To compensate for increased stress, regulatory mechanisms (such as osmoregulation) come at a significant energetic cost (Rivera-Ingraham and Lignot 2017). In turn, this can lead to an elevated oxygen demand and increased respiration rates (Rivera-Ingraham and Lignot 2017).

Previous experimental evidence suggests that impacts caused by decreased salinities would be lower for invasive species (Lenz et al. 2011). While *S. clava* had a lower respiration rate than *C. intestinalis* at all salinities, surprisingly we found limited evidence to suggest that, within the range tested, hyposalinity had a greater negative impact on *C. intestinalis*. Furthermore, given that previous evidence suggests the salinity tolerance range of *C. intestinalis* to be greater than *S. clava* (Dybern 1967; Lützen 1999; Carver et al. 2006; Davis et al. 2007), it is interesting to note that a behavioural response (unresponsive to stimuli) was observed in *C. intestinalis* at higher salinities (lower stress). Within this study, we did not measure recovery, ultimately this could be a determining factor when analysing the tolerance limits of either species.

Tolerance ranges can often vary among populations of the same species. Within the Baltic Sea, Dybern (1967) noted the high tolerance of *C. intestinalis* to low salinities of 11 in regions that are regularly subjected to these regimes. Similarly, Lützen (1999) noted that populations of *S. clava* within Danish fjords are able to survive regular periodic drops below salinities of 20, tolerating salinities documented to be lethal elsewhere. The individuals collected in this study were taken from a marina which was in close proximity to the Penfeld River in Brest. This could facilitate regular

exposure to low salinities, especially after strong rainfall. Over generations, species have the ability to adapt to new locations and conditions through natural selection and adaptive evolution (Colautti and Lau 2015). This adaptive ability seen in invasive species makes them particularly problematic within the marine environment. To fully appreciate the impact of invasive species, studies need to be conducted on the resistance of species to environmental pressures within and among their native and invasive ranges.

Conclusion

We found that the invasive S. clava displayed greater tolerance compared with C. intestinalis to both increased temperatures and to decreased salinities; however, limited differences in metabolic response between the two species were observed. This study is among the first to experimentally suggest upper limits on survival and metabolic response for either species to these common environmental stressors. This was due to limited prior experimental evidence for the newly taxonomically re-evaluated C. intestinalis and the invasive S. clava. As such, this study offers insights into the mechanisms behind the successful ability shown in these species to become invasive worldwide. Human-mediated dispersal of organisms is likely to select for the most tolerant, fast-growing and adaptive species. These species must be able to survive transport and subsequently become established within an area to truly become invasive. Future threats from climate change will only exacerbate the spread of invasive species. The wide tolerance ranges displayed here by S. *clava* are typical of invasive species worldwide, and as such, these factors must be taken into consideration to understand how and where species invasions are likely to occur.

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Compliance with ethical standards

Conflict of interest The authors declare no conflicts of interest.

Ethical approval All applicable international, national, and/or institutional guidelines for the care and use of animals were followed.

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