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The effect of short-term temperature exposure on vital physiological processes of mixoplankton and protozooplankton

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ABSTRACT

Sudden environmental changes like marine heatwaves will become more intense and frequent in the future. Understanding the physiological responses of mixoplankton and protozooplankton, key members of marine food webs, to temperature is crucial. Here, we studied two dinoflagellates (one protozo- and one mixoplanktonic), two ciliates (one protozo- and one mixoplanktonic), and two cryptophytes. We report the acute (24 h) responses on growth and grazing to a range of temperatures (5–34 °C). We also determined respiration and photosynthetic rates for the four grazers within 6 °C of warming. The thermal performance curves showed that, in general, ciliates have higher optimal temperatures than dinoflagellates and that protozooplankton is better adapted to warming than mixoplankton. Our results confirmed that warmer temperatures decrease the cellular volumes of all species. Q10 coefficients suggest that grazing is the rate that increases the most in response to temperature in protozooplankton. Yet, in mixoplankton, grazing decreased in warmer temperatures, whereas photosynthesis increased. Therefore, we suggest that the Metabolic Theory of Ecology should reassess mixoplankton’s position for the correct parameterisation of future climate change models. Future studies should also address the multigenerational response to temperature changes, to confirm whether mixoplankton become more phototrophic than phagotrophic in a warming scenario after adaptation.

1. Introduction

Climate change is expected to affect marine ecosystems and their biodiversity profoundly. One of the major issues surrounding this topic is the prediction that sea surface temperatures will rise continually (e.g., Xiao et al., 2019). In this regard, higher temperatures are predicted to increase both autotrophic and heterotrophic processes (such as photosynthesis and ingestion, respectively), albeit at different rates (e.g., Regaudie-de-Gioux and Duarte, 2012). In particular, the Metabolic Theory of Ecology (MTE – Brown et al., 2004) predicts that the activation energy (Ea) for the rate-limiting biochemical reactions of photosynthesis is significantly lower than the value for heterotrophic activities such as respiration and grazing (Regaudie-de-Gioux and Duarte, 2012; Rose and Caron, 2007). Therefore, in a warming scenario, marine ecosystems are expected to become less efficient at capturing carbon from the atmosphere, due to the prevalence of heterotrophic processes over autotrophic ones, at least over short time scales (Barton et al., 2020).

Still, this prediction is based on organisms that expressed either exclusive heterotrophy or autotrophy and did not consider photo-phagomixotrophy. The term mixoplankton has been recently coined to designate photo-phagotrophic protists (Flynn et al., 2019) and will be used throughout this work (exclusive phagoheterotrophs will be referred to as protozooplankton, and exclusive photo-autotrophs as phytoplankton). If we apply the MTE prediction as a universal law, we could also assume that, in mixoplankton, heterotrophic processes are expected to increase faster than autotrophic ones in response to warmer temperatures. This would shift the balance of photo/phagotrophy towards the latter mode of nutrition in mixoplankton.

It is important to mention that, irrespective of the trophic mode of nutrition, a change in metabolic rates due to temperature is going to repercuss on the performance of the individual, and thence on the population, and ultimately, on the entire ecosystem (Hochachka and
Somo, 2002). Furthermore, the temperature may also directly affect vertical and/or latitudinal distributions of organisms (An Gilletta Jr and Angilletta, 2009), which could be a source of unexpected biological interactions through predation or competition and affect the species composition of a given ecosystem (Montagnes et al., 2008). In addition, a direct consequence of climate change (yet understudied) is an increased frequency and intensity of extreme short-term events (Sal les et al., 2016) such as marine heatwaves, which are responsible for a sudden temperature change (Oliver et al., 2019). Experimenting on the ecosystem is unrealistic due to time and scale constraints because changes require longer periods of time to be noticeable, and sample representability is hard to determine. Thus, the best way to predict how will the ecosystem respond to a given stressor (such as temperature) is to experiment on a low organisational level, whose changes are likely going to affect the ecosystem (Zemos et al., 2010).

Therefore, measuring key metabolic rates on a diverse group of organisms within the ecosystem seems to be the ideal approach to study large ecosystem processes like the effects of temperature shifts. The visualisation of these effects requires the assembling of a thermal performance curve (An Gilletta Jr and Angilletta, 2009; Schulte et al., 2011). These curves are extremely useful in predicting the responses of populations to climate change, namely due to their possible incorporation into mechanistic models (Angert et al., 2011). Given the recent acknowledgement of the widespread occurrence of mixoplankton in microbial food webs, determining their thermal performance curves may promote an accurate integration into biogeochemical models (Mitra et al., 2014). Nevertheless, at the moment, data on the effects of temperature on key physiological parameters of mixoplankton are relatively scarce and contradictory (Cabrerozio et al., 2019; González-Ollala et al., 2019; Princtotta et al., 2016; Wilken et al., 2013).

Therefore, in this study, we aimed to determine the short-term (ca. 24 h) thermal performance curves for some phyto-, protozoa-, and mixoplanktonic species. We also measured respiration and photosynthetic rates on the phagotrophic species over a shorter temperature range to understand how temperature changes affect the internal metabolism of protozoo- and mixoplanktonic species. Altogether, the data collected with our experiments may, in future, fuel climate change models while attempting to clarify the place of mixoplankton within the MTE (either as enhanced producers or consumers in the ecosystem i.e., increased autotrophy or phagotrophy).

2. Methods

2.1. Cultures

We conducted the experiments with two protozooplankton species: the dinoflagellate Gyrodinium dominans (strain ICM-ZOO-GD001), and the ciliate Strombidium arenicola (strain ICM-ZOO-SA001). We also used two mixoplanktonic species from distinct functional groups (Mitra et al., 2016): the constitutive dinoflagellate Karldinium armiger (strain ICM-ZOO-KA001), and the plastidic-specialist ciliate Mesodinium rubrum (strain DK-2009). The two dinoflagellates and the ciliate S. arenicola were fed the cryptophyte Rhodomonas salina ad libitum during the up-scale period. To avoid the depletion of R. salina in the predator’s cultures, we supplied them with fresh cryptophytes every second or third day, depending on the predator species. M. rubrum was offered the cryptophyte Teleaulax amphioxia (strain K-1837) as prey in a proportion of ca. 1:5 (Smith and Hansen, 2007) during the up-scaling process. Protozooplankton and mixoplankton were maintained at ca. 100 μmol photons m−2 s−1, in autoclaved 0.1-μm filtered seawater. Both cryptophytes were grown in 1/2 medium (Guillard, 1975), prepared using autoclaved 0.1-μm filtered seawater, and irradiated at ca. 150 μmol photons m−2 s−1 provided by cool white fluorescent lights. To maintain the organisms under exponential growth (and within target concentrations), the stock cultures were diluted every 1–2 days with fresh medium (between 20 and 50% of the total volume). All cultures were kept at a salinity of 38 in a temperature-controlled room at 19 °C with a 10:14 L/D cycle.

2.2. Cell counts and volumes

Except for M. rubrum and its prey, all the organisms were counted and sized with a Beckman Coulter Multisizer III particle counter. M. rubrum may escape the current flow generated by the particle counter due to their sensitivity to shear and fast jump responses (Ferreira and Calbet, 2020). Therefore, cell counts of this ciliate using this instrument are often not representative of the concentration of the entire population. Accordingly, aliquots of M. rubrum, fixed in acidic Lugol’s solution (final concentration 2%), were prepared for all the treatments. A minimum of 300 predator and prey cells were counted using a Sedgewick-Rafter counting chamber. Additionally, 30 organisms were sized per replicate using the Fiji software (Schindelin et al., 2012); i.e., 90 cells were measured per temperature for the feeding and respiration experiments. Organismal volumes were estimated from linear dimensions using the following geometric shapes: for M. rubrum, a rotational ellipsoid, and for T. amphioxia, the added volume of a hemisphere and a cone (Smith and Hansen, 2007).

Since we noticed that M. rubrum and T. amphioxia cells enlarged when fixed in Lugol’s solution (using the previously described geometrical models), we conducted an independent trial where we sampled a single population of each species (for M. rubrum, n > 1.0 × 10³ cells; for T. amphioxia, n > 1.3 × 10⁶ cells) and ran an aliquot through the Beckman Coulter Multisizer III while fixing another in acidic Lugol’s (final concentration 2%). Despite not rendering trustable cell counts for M. rubrum, the electronic particle counter provided accurate volume estimations. We measured 200 organisms of each species from the fixed sample and obtained a conversion factor to correct the Lugol’s-preserved volumes (μm³) into live volumes (μm³) using the organisms measured with the electronic particle counter. For M. rubrum, Live Volume = 0.336 × Lugol Volume + 1169.443; for T. amphioxia Live Volume = 0.456 × Lugol Volume – 1.229 (Figs. S1 and S2 of the Supplementary Information).

Irrespective of the species, the changes in cellular volume (∆Volume, μm³) were used to assess the effect of temperature and were calculated according to Equation 1

\[ \text{Volume}_{\text{Temp}} = Vf_{\text{Temp}} - Vf_{\text{Temp}} \]

where Vf corresponds to the average volume of a cell (μm³) within a population after being exposed to a given target temperature (Temp, °C) for ca. 24 h. Vf, on the other hand, depicts the average volume of a cell (μm³) before exposure to the target temperature. Carbon values for all species were obtained from the pg Cμm³ ratio provided by Traboni et al. (2020) and used to determine C-specific rates.

2.3. Thermal performance curves

To assess the acute effects of temperature on the growth and grazing rates of protozooplanktonic and mixoplanktonic grazers, we exposed them to a wide range of temperatures (5–34 °C) for ca. 24 h with a 10:14 L/D regime at 100 μmol photons m−2 s−1. These temperatures were reached and maintained using recirculating water baths connected to individual aquarium chillers and heaters (TECO®). The incubations were conducted in triplicate experimental (predator and prey) and control (only prey) 132 mL Pyrex bottles. The bottles were submerged in the water baths during the incubation, and the temperature was monitored continuously using an Onset HOBO data logger. The target temperature was always reached within 20 min (from the initial stock temperature of 19 °C). Thermal performance curves of the growth rates of the cryptophytes R. salina and T. amphioxia, used as prey in the incubations, were obtained from the control bottles in the grazing experiments.
The experimental and control suspensions were prepared with the addition of 100 mL of fresh I/2 medium per L of suspension (final nutrient concentration equivalent to I/20 medium – Guillard, 1975). The objective was to avoid nutrient limitation for the prey during the incubation. During these experiments, K. armiger, G. dominans, and S. arenicola were fed R. salina, whereas M. rubrum was fed T. amplihoexia. All the experiments were conducted at saturating food concentrations (Table 1) to minimise the effect of different food concentrations on the measured ingestion rates. Predator concentrations were adjusted to allow ca. 30% of the prey to be consumed during the incubation while maintaining saturating food conditions.

All bottles were filled gradually, in three or four steps, using the corresponding experimental and/or control suspension, which was carefully mixed in between fillings. Additional experimental and control bottles were sacrificed at the beginning of the incubations to obtain the initial concentrations of the organisms. Cell numbers and volumes were obtained as described before. Growth and grazing rates at each target temperature were calculated after ca. 24 h of incubation using Frost (1972) and Heinbokel (1978) equations. Ingestion rates were deemed significant (i.e., not 0) only when the control and experimental bottles’ prey growth rates differed significantly (two-tailed Student’s t-test, P < 0.05). The temperature at which a given rate was maximised was termed $T_{\text{opt}}$ and the range of temperatures where this rate’s performance met or exceeded 80% of the observed at $T_{\text{opt}}$ was defined as $T_{\text{breath}}$ (Schulte et al., 2011).

### 2.4. Oxygen consumption and production rates

In addition to the feeding experiments, we conducted parallel trials at three different temperatures to quantify the consumption and production rates of oxygen during light and dark conditions. The chosen temperatures included the one used for the maintenance of parental cultures (i.e., 19 °C, and 3 °C below and above it. We used optical oxygen sensors (OxygenDipping Probe DP-PSt3, PresensH) at the beginning and at the end of the incubations (that lasted ca. 24 h) to determine oxygen concentrations. These experiments were conducted in triplicate experimental and control bottles, under a regular diel light cycle (light bottles; i.e., with a 10:14 L/D regime, 100 μmol photons m$^{-2}$ s$^{-1}$) or wrapped in aluminium foil throughout the incubation (dark bottles; i.e., with no light through the whole incubation). The control bottles contained only 0.1 μm-filtered seawater, whereas the experimental ones also contained grazers at a known concentration. The prey concentrations in the predator stock cultures were adjusted to guarantee their depletion on the night before the experiment, and therefore ensure a good physiological condition of the predators while eliminating the possible artefacts that co-existing prey could induce (Almeda et al., 2011; Calbet et al., 2022). Nevertheless, the absence of prey was further confirmed before the beginning of the experiments with the aid of the electronic particle counter. Additionally, initial bottles were also prepared in triplicate to assess the initial oxygen concentrations, necessary to compute the oxygen consumption rates later.

The oxygen consumption rates under darkness ($O_{\text{dark}}$, μmol O$_2$ L$^{-1}$ h$^{-1}$) were obtained considering triplicate dark bottles (i.e., ca. 24 h of darkness) using Equation (2) to determine the oxygen consumption of each individual dark bottle

$$O_{\text{dark}} = \frac{(C_{\text{f}} - C_{\text{i}})}{V_{\text{prop}}} - \frac{(E_{\text{Exp}} - E_{\text{Ctr}})}{i_{\text{Exp}}}$$

(2)

where $C_{\text{f}}$ corresponds to the oxygen concentration (μmol O$_2$ L$^{-1}$) inside the control bottles and Exp to the same parameter inside experimental bottles, and the subindex f and i correspond to the final and initial values, respectively. The incubation time (h) for the experimental bottles is represented by $i_{\text{Exp}}$ and for the control bottles by $i_{\text{Ctr}}$. The horizontal bars above some parcels of the equation (e.g., $E_{\text{Exp}}$) indicate that the average of the three replicates should be used.

The oxygen consumption rates obtained using Equation (2) were converted into per capita rates by dividing $O_{\text{dark}}$ by the average cell concentration of grazers in each bottle. These concentrations were obtained using Frost (1972) and Heinbokel (1978) equations after measuring the initial and final concentration of organisms as described before. Finally, oxygen consumption rates (under darkness) per unit of carbon per hour (i.e., respiration rates, R, μmol O$_2$ pg C$^{-1}$ h$^{-1}$) were obtained from the division of the last value by the average C concentration (pg C L$^{-1}$) in the same bottle, which was calculated from the C:μ m$^2$ ratio provided by Traboni et al. (2020). Notice that the calculation of R as described yields a positive value even though it is a C loss for the organism.

We considered only the triplicate light bottles to calculate oxygen consumption/production rates during the light period ($O_{\text{light}}$, μmol O$_2$ L$^{-1}$ h$^{-1}$) for both mixoplanktonic and protozooplanktonic grazers. We assumed that the respiration rate R was the same in the dark and light bottles (Wielgat-Rychert et al., 2017) and considered that our experimental setup for the latter comprised 14 h of darkness (as per the L/D cycle of the culture room). Equation (3) was then applied

$$O_{\text{light}} = \frac{\left[\left(E_{\text{Exp}} - (D_i C_{\text{Exp}}) \times 1.28\right) - \left(E_{\text{Ctr}} - (D_i C_{\text{Ctr}}) \times 1.28\right)\right]}{i_{\text{Exp}}} \times i_{\text{Exp}}$$

(3)

where $E_{\text{Exp}}$, $C_{\text{Exp}}$, $E_{\text{Ctr}}$, and the letters f and i have the same meaning as in Equation (2). The horizontal bars above specific parcels, as in Equation (2), also indicate average values. R is the respiration rate (μmol O$_2$ pg C$^{-1}$ h$^{-1}$) as calculated before from the dark bottles, and $C_{\text{Exp}}$ is the average concentration of C in the experimental bottle (from the grazer), as calculated using Frost (1972) equations. Per capita and per unit of carbon values were obtained as described before.

For mixoplanktonic species, $O_{\text{dark}} = R$ and $O_{\text{light}} = P$ (photosynthetic rate), according to Wielgat-Rychert et al. (2017). For protozooplankton, $O_{\text{light}}$ resulted in negative values, i.e., oxygen consumption during the hours of light and, therefore, light bottles were considered replicates from the dark incubations, and their average was used to determine R in protozooplankton (i.e., $O_{\text{light}}$:$O_{\text{dark}} = R$).

For the C-specific respiration (C losses), we multiplied R by the average respiratory quotient (moles of carbon dioxide produced per mole of oxygen consumed) of 0.89 (Williams and del Giorgio, 2005). The exact opposite, i.e., the molar ratio of oxygen produced to fixed carbon dioxide via photosynthesis, is called the photosynthetic quotient. Likewise, we multiplied P by the average photosynthetic quotient of 1.28 (Wielgat-Rychert et al., 2017) to obtain C-specific photosynthetic rates.

### 2.5. Activation energies and Q$_{10}$ coefficients

Activation energies (E$_a$, given in eV) can be obtained from the slope of the linear regression between the natural logarithm of a given rate versus the inverse of the absolute temperature (given in Kelvin, K) multiplied by the Boltzmann’s constant (8.62 × 10$^{-5}$ eV K$^{-1}$) (Vaquer-Sunyer et al., 2010). This plot is commonly referred to as an

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**Table 1**

Summary of the prey and predator concentrations used for the assembling of the thermal performance curves for the predator species, the protozooplanktonic G. dominans and S. arenicola, and the mixoplanktonic K. armiger and M. rubrum. All concentrations were based on published functional responses.

<table>
<thead>
<tr>
<th>Species</th>
<th>Target concentration, Cells ml$^{-1}$</th>
<th>Functional Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>G. dominans</td>
<td>100,000</td>
<td>Calbet et al. (2012)</td>
</tr>
<tr>
<td>S. arenicola</td>
<td>100,000</td>
<td>Ferreira et al. (2021)</td>
</tr>
<tr>
<td>K. armiger</td>
<td>100,000</td>
<td>Berge et al. (2008)</td>
</tr>
<tr>
<td>M. rubrum</td>
<td>15,000</td>
<td>Smith and Hansen (2007)</td>
</tr>
</tbody>
</table>
Arrhenius plot. Therefore, each physiological rate yields an individual Arrhenius plot for each species. To make it easier for the reader, we decided to convert $E_a$ into $Q_{10}$ coefficients, which represent the fold-increase in a given rate within a 10 °C variation, using Equation (4) (Vaquer-Sunyer et al., 2010)

$$Q_{10} = e^{(10 \frac{E_a}{R T})^2}$$

(4)

where $R$ is the gas constant (8.314 J mol$^{-1}$ K$^{-1}$), and $T$ is the mean absolute temperature for the range over which $Q_{10}$ was measured (upper and lower thermal extremes excluded – e.g., Eppley, 1972). For this calculation, $E_a$ were expressed in J mol$^{-1}$ using a conversion factor of 96486.9 (Vaquer-Sunyer et al., 2010).

3. Results

3.1. Temperature effects on tolerance and performance

The C-specific thermal performance curves for all six species are shown in Fig. 1 (for cell-specific rates, the reader is referred to Fig. S3), and the respective $T_{opt}$ and $T_{breadth}$ are shown in Table 2. All species displayed a thermal performance curve characterised by a gradual increase up to $T_{opt}$ followed by a sharp decline for both C-specific growth and grazing rates.

The C-specific thermal performance curves for growth of the cryptophytes R. salina and T. amphioxeia are displayed in Fig. 1a and b, respectively. The former exhibited a higher $T_{opt}$ and $T_{breadth}$ than the latter and showed positive C-specific growth rate in a wider temperature range as well (ca. 5.3–31.1 °C for R. salina, as opposed to 7.3–25.9 °C for T. amphioxeia). Among all species studied (prey or predator), increase up to $T_{opt}$ followed by a sharp decline for both C-specific growth and grazing rates.

<table>
<thead>
<tr>
<th>Species</th>
<th>$T_{opt}$</th>
<th>$T_{breadth}$</th>
<th>$T_{opt}$</th>
<th>$T_{breadth}$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Growth</td>
<td>Growth</td>
<td>Grazing</td>
<td>Grazing</td>
</tr>
<tr>
<td>Rhodomonas salina</td>
<td>25.07</td>
<td>10.61</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Teleaulax amphioxeia</td>
<td>21.90</td>
<td>5.99</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Gyrodinium dominans</td>
<td>25.74</td>
<td>8.63</td>
<td>22.62</td>
<td>7.67</td>
</tr>
<tr>
<td>Strombidium arenicola</td>
<td>32.52</td>
<td>11.05</td>
<td>30.49</td>
<td>9.71</td>
</tr>
<tr>
<td>Karlodinium armiger</td>
<td>16.01</td>
<td>11.39</td>
<td>16.01</td>
<td>14.65</td>
</tr>
<tr>
<td>Mesodinium rubrum</td>
<td>21.90</td>
<td>7.22</td>
<td>19.08</td>
<td>6.06</td>
</tr>
</tbody>
</table>

Table 2 Summary of $T_{opt}$ and $T_{breadth}$ (°C) for each species obtained from the species-specific thermal performance curves displayed in Fig. 1. NA = not applicable.

Fig. 1. C-specific thermal performance curves for the studied protists in terms of growth (inverted blue triangles) and ingestion (yellow circles): a and b) the phytoplankton R. salina and T. amphioxeia; c and d) the protozooplankton G. dominans and S. arenicola; e and f) the mixoplankton K. armiger and M. rubrum. The shaded areas limit the $T_{breadth}$ for each rate. M. rubrum exhibited non-significant ingestion rates at some temperatures (two-tailed Student’s t-test, $P > 0.05$). The negative ingestion rates in G. dominans were significant (two-tailed Student’s t-test, $P < 0.05$) and, therefore, shown. Error bars ± se.
T. amphioxea was the species with the narrowest $T_{\text{opt}}$ (Table 2). Indeed, this cryptophyte could only grow at rates up to 80% of the maximum rate between 16.3 and 22.3 °C.

The thermal performance curves of the two protozooplanktons can be found in Fig. 1c (G. dominans) and 1d (S. arenicola). These organisms tolerated wide variations in temperature, as seen by the range of temperatures exhibiting positive growth rates. For the dinoflagellate G. dominans the survival range broadened over ca. 23.6 °C difference (Fig. 1c). G. dominans was the only species showing significantly negative ingestion rates ($P < 0.05$ at ca. 5.6 °C), which were paired with negative growth rates. For the ciliate S. arenicola, positive growth rates were detected on all temperatures tested (5.6–34.4 °C – Fig. 1d). In addition, S. arenicola also showed the highest $T_{\text{opt}}$ for both rates across all species (ca. 32.5 °C and 30.5 °C for growth and grazing, respectively – Table 2).

Regarding mixoplankton (Fig. 1e and f), K. armiger showed a wide thermal performance curve, with the characteristic sharp decrease occurring ca. 9 °C above the $T_{\text{opt}}$, whereas for the remaining species it occurred always within 2 °C. The widening of K. armiger’s curve resulted in the largest $T_{\text{breadth}}$ among all species, both in terms of growth and grazing rates (Table 2). Conversely, among the four predators, M. rubrum was the one displaying the narrowest $T_{\text{breadth}}$, both in terms of growth and grazing rates (Table 2). In addition, the mixoplanktonic ciliate was the most sensitive species in our study, as its survivability range was narrower than in any other species (ca. 16.5 °C, between 7.3 and 23.8 °C). Finally, K. armiger showed the lowest $T_{\text{opt}}$ and was the only species whose $T_{\text{opt}}$ was lower than the maintenance temperature (ca. 19 °C) to which all species were exposed before the experiment.

### 3.2. Temperature effects on cellular volumes

The thermal performance curves also enabled the assessment of the overall effect of temperature on the $\Delta \text{Volume}$ (see the Methods section for the calculation procedure) of each target species (Fig. 2). All species showed a significant decrease in cellular volume ($P < 0.01$ in all instances) at higher temperatures, being this effect more evident in the predators than in the cryptophytes, as noticed by the higher correlation coefficients.

We also applied simple linear regression models between $\Delta \text{Volume}$ and C-specific growth rates (Fig. 3), and between C-specific ingestion rates and $\Delta \text{Volume}$ (Fig. 4). For R. salina the regression between $\Delta \text{Volume}$ and C-specific growth was not significant ($P = 0.09$ – Fig. 3a). Conversely, T. amphioxea displayed a significantly negative regression between $\Delta \text{Volume}$ and C-specific growth rates (Fig. 3b). The same pattern was observed in the ciliate S. arenicola (Fig. 3d), with the addition that $\Delta \text{Volume}$ was also negatively associated with ingestion rates (Fig. 4b). For both G. dominans and M. rubrum, the variation in volume was positively correlated with growth (Fig. 3c,f – $P < 0.01$); however, ingestion rates could not explain the variation in volume for both species (Fig. 4a,d – $P > 0.05$). Finally, K. armiger exhibited a unique pattern: the $\Delta \text{Volume}$ explained ca. 63% of the observed changes in C-specific ingestion (Fig. 4c).

### 3.3. Temperature effects on physiological rates

C-specific respiration rates of both protozooplanktonic grazers showed a significant increase ($P < 0.01$) in respiratory rates as temperature rose (Fig. 5a and b), whereas mixoplanktonic grazers seemed unaffected (Fig. 5c and d). For the ciliate M. rubrum, photosynthesis was also unaffected by temperature ($P > 0.05$ – Fig. 5d). However, K. armiger nearly doubled its C-specific photosynthetic rates (from ca. 0.41 d$^{-1}$ to ca. 0.78 d$^{-1}$, $P < 0.01$ – Fig. 5e) from 16.2 to 21.9 °C. Ciliates showed higher C-specific rates than dinoflagellates within a given trophic mode of nutrition. Altogether, this information resulted in distinct overall responses to temperature, as summarized by the rate-specific $Q_{10}$ coefficients (Table 3). The respective Arrhenius plots and $E_a$ can be found in Fig. S4 and Table S1 respectively.

The $Q_{10}$ coefficients calculated for growth and grazing rates were consistently higher in protozooplankton than in mixoplankton, irrespective of the species. In fact, growth and grazing rates displayed a $Q_{10} < 1$ for both mixoplanktonic predators, as opposed to an average $Q_{10}$ of 1.68 and 2.88 for growth and grazing, respectively, in the protozooplanktonic grazers. In addition, the $Q_{10}$ coefficients for grazing were the ones with the highest difference between trophic modes, being ca. 5.6 times higher in protozooplankton than in mixoplankton. Conversely, photosynthesis was the physiological process that varied the most in mixoplankton in response to temperature changes, with K. armiger exhibiting the highest fold increase (ca. 1.91 vs ca. 1.05 in M. rubrum). Regarding respiration rates, the two protozooplanktoners displayed a higher sensitivity to temperature than their mixoplanktonic counterparts, by exhibiting an average $Q_{10}$ of 1.76, compared to 1.15 in mixoplankton.

### 4. Discussion

The objective of this research was to assess the short-term physiological response of protozoa to a sudden variation in temperature. In this regard, our study evidenced that several key physiological parameters
are heavily modulated by temperature (Figs. 1 and 5), with grazing and photosynthesis being the highest temperature-dependent parameters in protozooplankton and mixoplankton, respectively (Table 3). In addition, we observed that higher temperatures implied smaller organisms (Fig. 2). The lack of an acclimation period is not a very common approach in studies aiming to determine the physiological consequences of temperature changes (Kang et al., 2020; Lim et al., 2019; Ok et al., 2019). Still, there are studies where metabolic responses have been directly measured after an acclimation period of 15 min (e.g., Padfield et al., 2016). The rationale behind our experimental design was to improve the similarities between our laboratory experiment and an extreme short-term temperature event in the field, such as marine heatwaves (whose frequency and intensity are projected to increase in the future – (Oliver et al., 2019), water mass displacements, etc. Therefore, our data intend to address a specific question, and we must highlight that different time scales allow different processes to occur, which could imply different effects. Indeed, Franzé and Menden-Deuer (2020) state that physiological acclimation can take up to 2.5 d °C⁻¹ when transitioning towards lower temperatures and 1.25 d °C⁻¹ when temperatures increase. Thus, our experiments may have slightly overestimated the variations of the measured rates in response to temperature (compared to more extensive time-scale studies). However, these differences were likely minor (as per the differences in non-acclimated vs acclimated populations (Franzé and Menden-Deuer, 2020; pers. obs.). Moreover, given the fast generation times of the species studied (Fig. S3), only ≤24 h experiments can capture the individual physiological response to temperature.

4.1. Temperature effects on thermal tolerance and performance

*K. armiger* showed a particularly wide T\text{breadth} in growth and ingestion rates (Fig. 1e and Table 2). It is not the first time a similar response is seen, as exemplified by its congener *Karlodinium veneficum* (Lin et al., 2018; Vidyarathna et al., 2020) and by other mixoplanktonic dinoflagellates (Kang et al., 2020; Lim et al., 2019; Ok et al., 2019). These wide T\text{breadth} could be the reason why these dinoflagellates possess a global distribution (Leles et al., 2019), as it would provide them with the necessary traits to colonise different environments characterised by variable temperatures (Angilletta Jr and Angilletta, 2009). The opposite (i.e., a narrow T\text{breadth}) was found in the ciliate *M. rubrum* and the cryptophyte *T. amphioxia* (Fig. 1b,f and Table 2). In the work of Fiorendino et al. (2020) the T\text{breadth} for these two species were similar to those obtained in our study; however, T\text{opts} were slightly higher than ours even though the strains used were the same. Still, in the work of Fiorendino et al. (2020) both species were adapted to a slightly higher temperature than in our experiments and they were acclimated for 2 days for every °C of variation until the experimental temperature was reached. Thus, this procedure may have increased their tolerance and performance to higher temperatures (Chakravarti et al., 2017), Gaillard et al. (2020) found similar growth performances for *T. amphioxia* in response to temperature. Thus, the combined assessment of our and other studies suggests that *M. rubrum* and its prey are tightly coupled in terms of thermal tolerance.

In addition, we have also confirmed that for the species investigated, protozooplanktonic predators are better adapted to a sudden increase of temperature than their mixoplanktonic counterparts, as seen by the higher average T\text{opts} (both in growth and ingestion rates) in the former group (Table 2). Regarding protozooplankton, we must highlight the negative ingestion and growth rates obtained at the lowest temperature in *G. dominans’* performance curve (Fig. 1c). These results denote a higher growth of the prey in the presence of the dinoflagellate than when incubated alone (controls), and likely resulted from an increase in the nutrient pool because of the death of grazers (e.g., Ferreira and Calbet, 2020). We attempted to eliminate this possible artefact by adding nutrients to the experimental suspensions; however, ammonium and urea, for example, can be released by dead microplankton (Caperon et al., 1979; Gao et al., 2018) and may explain the increased growth of *R. salina*. Franzé and Menden-Deuer (2020) previously reported mortality at similar temperatures, which could suggest that there is a threshold temperature for *G. dominans* around 5–6 °C. Finally, our results also indicate that the *R. salina* strain used in this study was better adapted to varying water temperatures than the one studied by Hammer et al. (2002), as seen by the better performance displayed at all temperatures.

4.2. Temperature effects on cellular volumes

Volume reductions due to temperature increases have been observed previously (Franzé and Menden-Deuer, 2020; Montagnes et al., 2008), and it has been proposed as a universal ecological response to increasing ambient temperatures (Daufresne et al., 2009; Sheridan and Bickford, 2011). Reductions in cell volume could be a consequence of individual cell shrinkage or higher cellular division rates. In addition, in the case of the grazers, changes in volume can also be a consequence of the ingestion of prey. Volume reductions have also physiological consequences. For instance, nutrient acquisition in phototrophs depends on the cellular surface/volume relationship (Pasciak and Gavis, 1974). Likewise, ingestion rates for planktonic grazers depend heavily on prey encounter rates, which is also a function of cell size (Kierboe and MacKenzie, 1995). Our study demonstrated that the ciliate *S. arenicola* exhibited a significantly negative slope of the linear regressions between C-specific ingestion and ΔVolume, and ΔVolume and C-specific growth (Figs. 3d
Therefore, it means that i) smaller *S. arenicola* grew faster than larger ones and that ii) smaller cells have higher C-specific ingestion rates. Being a protozooplanktonic grazer, the principal mechanism of C acquisition is through the ingestion of particulate matter. As such, we can conclude that the overall decrease in volume at higher temperatures (Fig. 2d) results from an enhanced cellular division rate, which in turn can only be attained due to higher C-specific ingestion rates. Similarly, *T. amphioxeia* also became smaller due to faster growth rates at higher temperatures. The conclusions for *S. arenicola* and *T. amphioxeia* are supported by the direct calculation of doubling times as \( \ln(2)/\mu \) (cell-specific) for the chosen temperatures (Figs. S3b and d).

On the contrary, species like the dinoflagellate *G. dominans* and the ciliate *M. rubrum* showed a positive regression between \( \Delta \text{Volume} \) and C-specific growth while displaying a non-significant relationship with C-specific ingestion rates (Fig. 3c,d and 4a,d). Thus, it seems that the smaller cells, observed at higher temperatures (Fig. 2c,f), are due to somatic reasons, i.e., higher temperatures shrink individual cells, although not necessarily as a consequence of higher cell-specific growth rates (Figs. S3c and f). On the other hand, *K. armiger* showed significant positive slopes between \( \Delta \text{Volume} \) vs C-specific growth and C-specific ingestion vs \( \Delta \text{Volume} \) (Figs. 3e and 4c). This means that the variation in volume (not cellular division – see Fig. S3e) can explain changes in growth, and that ingestion is the cause for the enlargement of the predator’s cell. Accordingly, at lower temperatures, the only logical conclusion is that *K. armiger* did not digest the ingested cells and did not divide, resulting in a very significant increase in its size (Fig. S5). Hence, we can conclude that the pattern seen in Fig. 2e was a consequence of the effect of temperature on grazing and not directly on *K. armiger*’s volume.

### 4.3. Temperature effects on physiological rates

According to our \( Q_{10} \) coefficients, only protozooplanktonic species are expected to increase their grazing rates in a sudden warming scenario, being the ciliate *S. arenicola* the species benefiting the most, with a \( Q_{10} \) of 3.09 (Table 3). In the case of mixoplankton, both species exhibited a value < 1, which indicates that an increase in the ambient temperature will cause a decreased ingestion of particulate food. The magnitude of the effects of temperature on the four grazers agrees with the maximum ingestion rates of their respective functional responses. Indeed, *S. arenicola* consumed as much as 120 *R. salina* predator\(^{-1}\) d\(^{-1}\) (Ferreira et al., 2021), whereas *M. rubrum* only ate ca. 5 *T. amphioxeia* predator\(^{-1}\) d\(^{-1}\) (Smith and Hansen, 2007). *G. dominans* and *K. armiger* stand in between the two ciliates, with the protozooplankter eating ca. 20 *R. salina* predator\(^{-1}\) d\(^{-1}\) (Calbet et al., 2013) and the mixoplankter ca. 10 *R. salina* predator\(^{-1}\) d\(^{-1}\) (Berge et al., 2008). This correlation could be a consequence of the C requirements for each species and their mechanism of acquisition (i.e., how much phagotrophy contributes to the overall C budget).

Regarding respiration rates, the effect of temperature was also higher in protozooplankton than in mixoplankton, as seen by the higher \( Q_{10} \) for this rate in the first group. This result implies that protozooplankters will lose proportionally more C through respiration in a warming scenario than mixoplankters. This is likely a consequence of internal photosynthetic mechanisms in mixoplankton, which often (if not always) prioritise internal over external C sources i.e., the internal recycling of C decreases their overall void of C (Flynn and Mitra, 2009). Conversely, for protozooplankton, grazing is the only source of C acquisition and, therefore, it seems logical to find a correlation between C intake by
feeding and C losses due to respiration (as both parameters are integrated into an organism’s C budget). It has been reported that residual photosynthesis may occur in protozooplankton due to the presence of algae food in vacuoles (Ferreira et al., 2021; Tarangkoon and Hansen, 2011). Still, in our experiment, all predators were allowed to deplete their co-existent prey on the night before the experiment. Thus, even though photosynthesis was not ruled out as a hypothetical activity in protozooplankton, the lack of prey in the respiration/photosynthesis experiment minimised this potential problem. In addition, respiration rates are typically higher in the presence of food than in its absence (Calbet et al., 2022). In any case, O_{light} for protozooplankton resulted in negative values, i.e., oxygen consumption during the hours of light. These results were typically slightly lower (on average ca. 5.3% lower) than those in the dark incubations (O_{dark}), although differences were never statistically significant (Student’s t-test, P > 0.05 on all cases) and, therefore, dark and light bottles were considered as replicates for the measurement of respiration rates in protozooplankton.

One interesting outcome of our experiments comes from the analysis of photosynthesis in both K. armiger and M. rubrum, in particular, in light of the Q_{10} coefficients for growth. It seems that the ciliate, whose mode of nutrition is primarily autotrophic (Smith and Hansen, 2007), might not benefit much from a sudden increase of temperature (Q_{10} for photosynthesis ca. 1.11 – Table 3), as predicted by the MTE for autotrophic processes (Brown et al., 2004). On the other hand, K. armiger, increased its photosynthetic rate by ca. 1.91 times in less than 6 °C, which resulted in a very high Q_{10} for this process in the dinoflagellate (ca. 3.16 – Table 3). Nevertheless, these increased photosynthetic rates were incapable of sustaining the growth of both predators as temperature rises (Q_{10} for growth is < 1), which suggests that grazing plays an important part in the overall metabolism of these organisms (Q_{10} for grazing < 1 for both species). Unsurprisingly, a severely reduced grazing in M. rubrum (Q_{10} = 0.33) had a lower effect on growth (Q_{10} = 0.92) than the one demonstrated by K. armiger with a smaller reduction in grazing (Q_{10} for grazing = 0.88; Q_{10} for growth = 0.80). This is likely entirely dependent on their specific reliance on auto/heterotrophic mechanisms of C acquisition, as the ciliate is primarily autotrophic and the dinoflagellate is a voracious feeder (Berge and Hansen, 2016; Smith and Hansen, 2007).

Considering all four physiological rates measured in this study, we can attempt to place these organisms within the MTE framework. A critical aspect that is conserved in both mixoplanktonic predators is that

Table 3

<table>
<thead>
<tr>
<th>Species</th>
<th>Growth</th>
<th>Grazing</th>
<th>Respiration</th>
<th>Photosynthesis</th>
</tr>
</thead>
<tbody>
<tr>
<td>G. dominans</td>
<td>2.01</td>
<td>2.66</td>
<td>1.67</td>
<td>NA</td>
</tr>
<tr>
<td>S. arenicola</td>
<td>1.34</td>
<td>3.09</td>
<td>1.85</td>
<td>NA</td>
</tr>
<tr>
<td>K. armiger</td>
<td>0.80</td>
<td>0.88</td>
<td>1.19</td>
<td>3.16</td>
</tr>
<tr>
<td>M. rubrum</td>
<td>0.92</td>
<td>0.33</td>
<td>1.10</td>
<td>1.11</td>
</tr>
</tbody>
</table>

Q_{10} for every physiological rate ascertained in this study for the predator species. Q_{10} was calculated using Equation (4) (Vaquer-Sunyer et al., 2010). NA = not applicable.

Fig. 5. C-specific respiration (red diamonds) and photosynthetic (green squares) rates for a) G. dominans, b) S. arenicola, c) K. armiger, and d) M. rubrum for temperatures between 16.2 and 21.9 °C. Non-significant regressions (P > 0.05) are depicted with a dotted line, whereas significant regressions are displayed with a solid line (** implies P < 0.01).
photosynthesis is the rate that benefits the most from temperature (although the differences across rates are minor in *M. rubrum*). In addition, grazing was always hindered in a sudden warming scenario in both species (seen by a Q10 < 1 – Table 3). Moreover, digestion rates depend on the ambient temperature but vary similarly in mixoplanktonic and protozooplanktonic grazers (*Fenchel, 1975; Li et al., 2001*). This particular combination of factors suggests that both mixoplanktonic species (irrespective of their taxonomic group) increase their auto/heterotrophic ratio at higher temperatures, as opposed to the predictions of the MTE for strict autotrophic and heterotrophic organisms (*Brown et al., 2004* and to some experimental studies as well (*Cabreroz et al., 2019; Wilken et al., 2013*). Nevertheless, our results are not the first to report an atypical behaviour of mixoplankton in light of the MTE projections. For example, a direct measurement of the contribution of grazing to the total metabolic budget in the bacterivore mixoplankter *Dinobryon sociale* resulted in a higher contribution of photosynthesis at higher temperatures (*Princotta et al., 2016*). Similarly, *Gonzalez-Olalla et al. (2019)* assessed the effect of temperature on two bacterivores and concluded that warmer temperatures shifted the overall metabolism towards an increased photosynthesis in both species. Also, *Ok et al. (2019)* studied the mixoplanktonic dinoflagellate *Takayama helix* (same family as *K. armiger*) and noticed increased growth rates paired with insignificant changes in ingestion rates in a wide temperature range. *Lim et al. (2019)* and *Kang et al. (2020)* noticed the same pattern in the mixoplanktonic dinoflagellates *Alexandrium pohangense* and *Yihiella yeosuensis*, respectively. Altogether, the results from these latter three works hint at a possibly higher phototrophic contribution to the overall metabolism in these dinoflagellate species, although this variable was not directly measured in their study.

Still, our results do not question prior estimations of Ea in phototrophs and heterotrophs based on growth rates (e.g., *Rose and Caron, 2004*), although this variable was not directly measured in their study. Nevertheless, the recent evidence demonstrated that the Ea values for growth are widely variable among different taxonomic groups (*Chen and Laws, 2017*). In addition, the nutritional plasticity of mixoplankton has been pointed as a possible source of error between theoretical and observed Ea in microplankton (*Wang et al., 2019*). Therefore, as our results support an increased photosynthesis in mixoplankton at higher temperatures, we contribute to the body of literature that deviates mixoplankton from the MTE. This conclusion means that such a change in nutritional strategies will likely impact biogeochemical cycles and reinforces the need to integrate mixoplankton in current ecosystem models (*Wilken et al., 2018*).

### 4.4. Final remarks

It is important to stress the laboratory nature of this study and the inherent adaptation to a constant temperature in the species (and strains) considered. In spite of being a perfectly natural approach to study microplanktonic communities for logistical reasons (*Flynn et al., 2019*), field communities likely experience temperature oscillations within a day (e.g., *Olives et al., 2022*). These natural die variations may decrease the overall effect of temperature on the physiological rates assessed in our study since the effects of adaptation to a specific set of abiotic conditions is mostly absent in the field. Indeed, future studies should also address the multigenerational response to temperature changes since a general (and gradual) increase in the oceanic temperature is also expected due to climate change (*Xiao et al., 2019*). Accordingly, adaptation will likely be reflected in the biological rates and overall metabolism, meaning that these changes must also be incorporated in future modelling predictions (*Calbet and Saiz, 2022*). In this regard, evidence from evolutionary studies suggests that, despite having a stronger temperature dependence, heterotrophic processes are balanced with autotrophic ones with passing generations, which culminates with higher C fixation rates in a future warming scenario (*Barton et al., 2020; Padfield et al., 2016*). Nevertheless, the data presented in this work should assist in comprehending the effect of climate change in marine protistan communities regarding short-term temperature events such as marine heatwaves. Finally, our study contributes to the correct placement of mixoplankton within the MTE, which may be crucial for the accurate projection of climate change in the future.

### Author contributions

All authors conceptualized the experiments. G.D.F. and A.G. prepared the cultures and conducted the experiments. A.C. and E.S. provided material, facilities and assistance to the development of the experiments. G.D.F., A.C., and E.S. analysed the data. G.D.F. and A.C. prepared the original draft manuscript. All authors read, contributed, and approved the final version of the manuscript thus justifying all authorships.

### Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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### Appendix A. Supplementary data

Supplementary data to this article can be found online at [https://doi.org/10.1016/j.marenvres.2022.105693](https://doi.org/10.1016/j.marenvres.2022.105693).

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