1	Cold-induced [Ca ²⁺] _{cyt} elevations function to support osmoregulation in marine diatoms
2	
3	Friedrich H. Kleiner ^{1,2} , Katherine E. Helliwell ^{1,3} , Abdul Chrachri ¹ , Amanda Hopes ⁴ , Hannah
4	Parry-Wilson ^{1,2} , Trupti Gaikwad ¹ , Nova Mieszkowska ^{1,5} , Abdul Chrachri ¹ , Thomas Mock ⁴ ,
5	Glen L. Wheeler ¹ *, Colin Brownlee ¹ *
6	
7	
8	1 The Marine Biological Association of the United Kingdom, The Laboratory, Citadel Hill,
9	Plymouth, Devon, PL1 2PB, UK
10	2 School of Ocean and Earth Science, University of Southampton, Southampton, SO14 3ZH,
11	UK
12	3 Biosciences, College of Life and Environmental Sciences, University of Exeter, Exeter EX4
13	4QD, UK
14	4 School of Environmental Sciences, University of East Anglia, Norwich Research Park,
15	Norwich, UK
16	5 School of Environmental Sciences, University of Liverpool, Jane Herdman Building,
17	Liverpool, L69 3GP, UK
18	
19	
20	
21	*Corresponding authors: <u>glw@mba.ac.uk</u> , cbr@mba.ac.uk
22	

23 Abstract

24

25 Diatoms are a group of microalgae that are important primary producers in a range of open 26 ocean, freshwater and intertidal environments. The latter can experience significant long- and short-term variability in temperature, from seasonal variations to rapid temperature shifts 27 caused by tidal immersion and emersion. As temperature is a major determinant in the 28 distribution of diatom species, their temperature sensory and response mechanisms likely 29 have important roles in their ecological success. We have examined the mechanisms diatoms 30 use to sense rapid changes in temperature, such as those experienced in the intertidal zone. 31 We find that the diatoms *Phaeodactylum tricornutum* and *Thalassiosira pseudonana* exhibit a 32 transient cytosolic Ca^{2+} ($[Ca^{2+}]_{evt}$) elevation in response to rapid cooling, similar to those 33 observed in plant and animal cells. However, $[Ca^{2+}]_{cyt}$ elevations were not observed in 34 response to rapid warming. The kinetics and magnitude of cold-induced [Ca²⁺]_{cyt} elevations 35 correlate with the rate of temperature decrease. We do not find a role for the $[Ca^{2+}]_{cvt}$ 36 elevations in enhancing cold tolerance, but show that cold shock induces a Ca²⁺-dependent K⁺ 37 efflux and reduces mortality of P. tricornutum during a simultaneous hypo-osmotic shock. As 38 39 inter-tidal diatom species may routinely encounter simultaneous cold and hypo-osmotic shocks during tidal cycles, we propose that cold-induced Ca²⁺ signalling interacts with 40 osmotic signalling pathways to aid in the regulation of cell volume. Our findings provide 41 insight into the nature of temperature perception in diatoms and highlight that cross-talk 42 43 between signalling pathways may play an important role in their cellular responses to multiple 44 simultaneous stressors.

46 Introduction

Diatoms are a group of silicified unicellular algae that represent one of the most important primary producers in modern oceans. They are abundant in diverse marine environments, most notably in polar and temperate upwelling regions, where they play a critical role at the base of the marine food web (Malviya et al., 2016). Diatom communities are abundant across a broad temperature range in the surface ocean from sea ice to tropical oceans. Diatoms are also important primary producers in freshwater and brackish ecosystems, where they likely encounter an even greater range of temperatures (Souffreau et al., 2010).

Global rises in surface temperature due to anthropogenic CO_2 emissions are set to have 54 55 profound influence on marine ecosystems (Gattuso et al., 2015). These future changes in our 56 climate will also increase the variability of temperature regimes and the prevalence of extreme events, such as marine heatwaves, that may co-occur with other stressors such as low pH or 57 58 deoxygenation (Harley et al., 2006; Smale et al., 2019; Gruber et al., 2021). Understanding the physiological response of diatoms and other marine phytoplankton to changes in global 59 60 temperature regimes is therefore of the utmost importance. Temperature has an important impact on diatom cell physiology, influencing cell size and formation of the silica frustule 61 (Montagnes and Franklin, 2001; Svensson et al., 2014; Javaheri et al., 2015). Individual 62 63 species display a thermal niche with distinct temperature growth optima that reflect their 64 natural environment (Liang et al., 2019). The upper and lower thermal tolerance limits, rather 65 than the optima themselves, appear to have the greatest influence on the distribution of 66 individual diatom species (Anderson and Rynearson, 2020), with temperatures in excess of 67 the upper thermal tolerance limits leading to a rapid increase in the rates of cell death (Baker 68 and Geider, 2021).

69 Many of these studies have focussed on the physiological responses of diatoms to longer term 70 changes in temperature. However, diatoms will also experience short term temperature 71 variations within their natural habitat. This is particularly so for those species that inhabit 72 intertidal rocky shores or estuarine habitats where immersion and emersion is associated with 73 rapid and regular temperature fluctuations. Rapid temperature changes are potentially highly 74 damaging to diatom cells, demonstrated by their much greater vulnerability to abrupt rather 75 than gradual temperature increases (Souffreau et al., 2010). Temperature variability may also 76 have an important influence on the ability of diatoms to adapt to their thermal niche, as 77 Thalassiosira pseudonana exhibited accelerated adaptation to higher temperatures under a 78 fluctuating temperature regime (Schaum et al., 2018). Despite the importance of thermal

tolerance in diatom physiology and ecology, relatively little is known about the physiological
mechanisms that allow diatoms to perceive and respond to changes in temperature,
particularly during short-term fluctuations.

82 Many of the cellular mechanisms involved in temperature sensing in eukaryotes involve temperature-induced changes in the structure of nucleic acids, proteins or biological 83 membranes that lead to a range of downstream physiological responses (Sengupta and 84 Garrity, 2013). Ca²⁺-dependent signalling mechanisms play an important role in these 85 temperature sensing pathways. In animal cells, heat stress is associated with Ca²⁺ influx into 86 the cytosol via the TRPV family of temperature-sensitive ion channels (Xu et al., 2002; 87 Clapham and Miller, 2011). Ca²⁺ signalling also plays a role in sensing low temperature in 88 animals, for example underpinning the rapid cold hardening response of insects (Teets et al., 89 2013). Land plants also employ Ca²⁺ signalling mechanisms in their response to both low and 90 high temperatures. Rapid cooling of plants induces a transient cytosolic Ca^{2+} ($[Ca^{2+}]_{cyt}$) 91 elevation, which leads to changes in gene expression and the establishment of cold tolerance 92 (Knight et al., 1996; Tahtiharju et al., 1997; Knight and Knight, 2012). Some plants, such as 93 the moss *Physcomitrium*, also display $[Ca^{2+}]_{evt}$ elevations in response to heat shock (Saidi et 94 al., 2009). In other plants, such as Arabidopsis, high temperatures do not induce $[Ca^{2+}]_{cvt}$ 95 elevations, although Ca²⁺ elevations are observed within the chloroplast (Lenzoni and Knight, 96 2019). Potential temperature sensors in plants include the cold sensitive COLD1/RGA1 97 complex in *Oryza sativa*, which is proposed to either function as a Ca^{2+} channel or to activate 98 other Ca²⁺ channels (Ma et al., 2015). Specific cyclic nucleotide-gated ion channels and 99 annexins may also play a role in temperature sensing pathways, with mutant strains in 100 Physcomitrium, O. sativa and Arabidopsis exhibiting diminished [Ca²⁺]_{evt} elevations in 101 102 response to cold- and heat shock (Cui et al., 2020; Liu et al., 2021). However, it is currently unclear whether these ion channels sense temperature directly or are activated indirectly, e.g. 103 104 through changes in membrane rigidity (Plieth et al., 1999) or the cytoskeleton (Pokorna et al., 2004). 105

Our understanding of Ca^{2+} signalling in diatoms remains in its infancy, although Ca^{2+} dependent signalling mechanisms have been identified in response to a range of environmental stimuli, such as the supply of nutrients (phosphate and iron), hypo-osmotic shock and the detection of toxic aldehydes (Falciatore et al., 2000; Vardi et al., 2006; Helliwell et al., 2021; Helliwell et al., 2021). Initial experiments using *Phaeodactylum tricornutum* cells expressing the bioluminescent Ca^{2+} reporter aequorin did not detect $[Ca^{2+}]_{cyt}$

elevations in response to low (4 °C) or high (37 °C) temperature (Falciatore et al., 2000). More recently, genetically-encoded fluorescent Ca^{2+} reporters have been successfully expressed in *P. tricornutum* and *T. pseudonana*, enabling high resolution imaging of $[Ca^{2+}]_{cyt}$ elevations in single diatom cells (Helliwell et al., 2021; Helliwell et al., 2021). These advances will now allow detailed examination of diatom signalling in response to range of stimuli, including temperature.

In this study we set out to examine the ability of diatoms to sense short-term changes in 118 temperature. In particular, we examined whether the well-characterised $[Ca^{2+}]_{evt}$ elevations 119 observed in animal and plant cells in response to rapid changes in temperature were conserved 120 121 in diatoms. Using the model species P. tricornutum and T. pseudonana, which can both 122 inhabit coastal environments that experience variable temperature regimes (De Martino et al., 2007; Alverson et al., 2011), we found that diatoms consistently exhibit a $[Ca^{2+}]_{cvt}$ elevation 123 in response to cold shock, but do not exhibit [Ca²⁺]_{evt} elevations in response to elevated 124 temperature. We did not find a requirement for cold shock-induced Ca²⁺ signalling in 125 increasing tolerance to low temperatures, but found that cold shock increases resistance to 126 simultaneous hypo-osmotic shocks, suggesting that integration of multiple signalling inputs 127 128 may contribute to an enhanced ability to respond to these environmental stimuli.

131 Methods

132 <u>Recording of rockpool temperature</u>

Temperature data was recorded using a 27 mm Envlogger v2.4 (ElectricBlue, Porto, Portugal)
encased in acrylic resin, recording in 30 minute intervals with a resolution of 0.1 °C. The
Envlogger was secured to the substrate using Z-Spar A-788 epoxy resin roughly 3 cm below
the surface waters of a shallow midshore rockpool measuring approximately 8 cm deep at
Looe Hannfore, Cornwall, UK (50.3411, -4.4598) from 1/7/2019 to 7/7/2019.

138 Strains and culturing conditions

The wild type P. tricornutum strain used in this study was CCAP 1055/1 (Culture Collection 139 of Algae and Protozoa, SAMS, Scottish Marine Institute, Oban, UK). A P. tricornutum strain 140 transformed with the R-GECO1 Ca²⁺ biosensor (PtR1) and the three *eukcata1* knock-out 141 strains in this line (labelled A3, B3 and B6) were generated as described previously(Helliwell 142 143 et al., 2019). The T. pseudonana strain expressing the R-GECO1 biosensor (TpR1) was generated as described in Helliwell et al (2021). Cultures were maintained in natural seawater 144 with f/2 nutrients(J C Lewin and Guillard, 1963; Guillard, 1975); modified by the addition of 145 106 µM Na₂SiO₃,5H₂O and the exclusion of vitamins (P. tricornutum only). For imaging 146 147 experiments, cells were acclimated to an artificial seawater (ASW) medium for minimum 10 148 days prior to analysis. ASW contained 450 mM NaCl, 30 mM MgCl₂, 16 mM MgSO₄, 8 mM 149 KCl, 10 mM CaCl₂, 2 mM NaHCO₃, 97 µM H₃BO₃, f/2 supplements and 20 mM HEPES (pH 8.0). Cultures were grown at 18 °C with a 16:8 light/dark cycle under illumination of 50 µmol 150 $m^{-2} s^{-1}$. 151

152 Epifluorescence imaging of R-GECO1 fluorescence

153 $500 \ \mu L$ of cell culture was added to a 35 mm microscope dish with glass coverslip base (In 154 Vitro Scientific, Sunnyvale, CA, USA) coated with 0.01% poly-L-lysine (Merck Life Science 155 UK, Gillingham, Dorset) to promote cell adhesion to the glass surface. Cells were allowed to 156 settle for 5-20 minutes at room temperature (RT) under light. R-GECO1 was imaged using a 157 Leica DMi8 inverted microscope (Leica Microsystems, Milton Keynes, UK) with a 63x 1.4NA oil immersion objective, using a Lumencor SpectraX light source with a 541-551 nm 158 159 excitation filter and 565-605 nm emission filter. Images were captured with a Photometrics 160 Prime 95B sCMOS camera (Teledyne Photometrics, Birmingham, UK). Images were 161 captured at 3.33 frames per second using Leica application suite X-software v.3.3.0.

162 Administration of temperature shocks to cells in the imaging setup

The dish was perfused with ASW without f/2 nutrients at a standard flow rate of 16 mL min⁻¹. 163 To achieve rapid changes in temperature in the dish, the perfusion was switched between 164 165 solutions of different temperature to achieve target temperatures of approximately 10, 22 or 166 30 °C respectively. Actual dish temperature was recorded using a Firesting micro optical 167 temperature sensor (Pyroscience GmbH, Aachen, Germany). For the majority of experiments, cells were perfused with warmer media (dish temperature 30 °C) for 1 minute prior to 168 169 application of the cooling shock (these conditions reflect those observed in the rockpool 170 observations). The perfusion flow rate was altered to achieve different temperature change 171 rates. As cooling rate was not linear, the maximum cooling rate was defined as the largest 172 decrease temperature within a one second period.

173 Application of inhibitors and elicitors

External Ca^{2+} was removed by perfusion with ASW without $CaCl_2$ containing 200 μ M EGTA. Ruthenium red (RR) was added to cells at a final concentration of 10 μ M 5 minutes prior to cold shock treatment. Menthol was prepared as a 1 M stock solution in DMSO and used at concentration of 1 mM, resulting in a final DMSO concentration of 0.1%.

178 Processing of imaging data

179 Images were processed using LasX software (Leica). The mean fluorescence intensity within 180 a region of interest (ROI) over time was measured for each cell by drawing an ROI 181 encompassing the whole cell. Background fluorescence was subtracted from all cellular F 182 values. The change in the fluorescence intensity of R-GECO1 was then calculated by normalizing each frame by the initial value (F/F₀). [Ca²⁺]_{cyt} elevations were defined as any 183 increase in F/F_0 above a threshold value (1.5). The duration of a $[Ca^{2+}]_{cyt}$ elevation was defined 184 as the peak width at half maximal amplitude. To visualise the spatial distribution of a $[Ca^{2+}]_{cvt}$ 185 186 elevation, each frame was divided by a corresponding background image generated by 187 applying a rolling median (30 frames) to the image-series (Image J). The resultant time series 188 images were pseudo-coloured to indicate changes in fluorescence.

189 <u>Statistical analysis</u>

190 Graphs and statistical analyses were performed using Sigmaplot v14.0 (Systat Software,191 Slough, UK). Error bars represent standard error of the mean. Unless indicated otherwise,

imaging experiments were repeated three times with independent cultures on different days toensure reproducibility of the response.

Normal distribution of respective datasets was tested using Shapiro-Wilk's normality test.
When passed, statistical analysis of datasets with two groups were done with Student's t-test,
and when not passed with Mann-Whitney's rank sum test. Statistical analysis of datasets with
more than two groups were performed using an ANOVA followed by a Holm-Sidak post-hoc
test when the normality test was positive. When the normality test was negative, KruskalWallis' 1-way Analysis of Variance on Ranks was used instead. All statistical tests were
performed with Sigmaplot v14.0.3.192 (Systat software Inc).

201 <u>Growth at different temperatures after a cold shock.</u>

For the growth curves, cells were grown to mid exponential phase $(2.73 \times 10^6 \text{ cells mL}^{-1})$. The culture was divided into 10 mL aliquots and cells were pelleted by centrifugation (4000 x g at 18 °C). Cells were washed in 40 mL ASW +/- Ca²⁺ and pelleted again by centrifugation. Cells were then resuspended in their respective treatments (20 mL of ASW +/- Ca²⁺ at 18 °C or 4 °C to administer a rapid cold shock). After 10 minutes, 2 mL of each culture was used to inoculate culture vessels containing 18 mL of standard F/2 media (approx. starting density of 6.8×10^4 cells mL⁻¹) and cultures were grown at 18 °C or 4 °C.

209 <u>Cell survival during hypo-osmotic shock</u>

To examine the effect of temperature on cell viability during hypo-osmotic shock, 10 mL of a 210 late log phase culture (6 x 10^6 cells mL⁻¹) were pelleted by centrifugation (4000 x g at 18 °C). 211 Cells were washed twice with 10 mL ASW $+/-Ca^{2+}$ and 250 µL aliquots were taken. To apply 212 the hypo-osmotic and cold shock treatments, 750 μ L of ASW (+/- Ca²⁺) or deionised water at 213 two different temperatures (20 °C or 4 °C) were added to each tube. Addition of water results 214 215 in a severe hypo-osmotic osmotic shock (final concentration 25% ASW) simultaneously with 216 the temperature shock. The cells were then incubated at their respective temperatures for 3 217 minutes prior to addition of 5 µM Sytox Green (Thermo Fisher Scientific, Loughborough, 218 UK). All treatments were then incubated at 20 °C 15 min in darkness. Cell viability was measured with a LUNA-FL[™] Dual Fluorescence Cell Counter (Logos Biosystems, Villeneuve 219 d'Ascq, France) to count live (displaying red chlorophyll fluorescence) versus dead cells 220 (Sytox Green stain) with following settings: Excitation intensity green = 11, red = 7, count 221 222 threshold for both = 3.

Quantification of K+ efflux in P. tricornutum populations using K+ -selective microelectrodes 223

 K^+ microelectrodes were fabricated as described previously (Helliwell et al., 2021). Clark 224 GC-1.5 borosilicate glass capillaries (Harvard Apparatus, Cambridge, UK) were pulled to a 225 fine point using a P-97 pipette puller (Sutter, Novato, CA). The pipette tips were then gently 226 broken to produce a diameter of ca 10-20 µm. The capillaries were silanised by exposure to 227 228 N,N-dimethyltrimethylsilylamine (TMSDMA) vapour at 200 °C for 20 minutes within a closed glass Petri dish. The K⁺ microelectrodes were prepared by introducing K⁺ ionophore I 229 (Sigma Aldrich, Gillingham, Dorset, UK) into the pipette tip by suction. Pipettes were then 230 231 back-filled with the filling solution (100 mM NaCl, 20 mM HEPES pH 7.2 and 10 mM 232 NaOH). The reference electrode was filled with 3 M KCl and data were recorded using an 233 AxoClamp 2B amplifier, with pClamp v10.6 software (Molecular Devices, CA, USA). Each 234 K⁺ microelectrode was calibrated using a two-point calibration with standard KCl solutions. The mean slope of the calibrated electrodes was 53.0 ± 1.3 mV per decade (\pm SE). 235 For the measurements, 10 mL of *P. tricornutum* cells from exponential culture containing 10^6 -236

10⁷ cells mL⁻¹ were centrifuged at 4000 rpm for 10 min and re-suspended in 1 mL of ASW. 237

The cells were then allowed to settle on a poly-L-lysine coated microscope dish. Cells were 238

perfused with ASW or ASW $-Ca^{2+}$ (0 μ M Ca^{2+} + 100 μ M EGTA) and cold shocks were 239

- applied as described for the microscopy observations. Control experiments were performed in the absence of P. tricornutum cells to ensure that the change in temperature did not alter the 241
- performance of the K⁺ microelectrodes. 242

243

245 **<u>Results</u>**

246 Rapid changes in temperature in inter-tidal environments

P. tricornutum was first isolated from a tidal pool in the UK and has since been identified in a 247 248 range of coastal and brackish habitats (De Martino et al., 2007). To assess the dynamic 249 temperature regimes potentially experienced by intertidal diatoms, we measured the 250 temperature of a tidal pool located on the upper region of a rocky shore (South Cornwall, UK) 251 over a 7 d period during July (UK summer). Temperatures within the pool were very stable 252 around 15°C during immersion at high tide (Fig 1). However, at low tides temperatures in the 253 exposed tidal pool rose significantly during the day (up to 30 °C) and decreased at night (to 254 12 °C), before being rapidly restored to the bulk seawater temperature by the immersion of 255 the pool at high tide. These data illustrate that diatoms inhabiting intertidal environments in 256 temperate regions will regularly experience periods of significant warming followed by rapid cooling. The fluctuations in temperature are likely to be even greater in smaller volumes of 257 258 water, such as the surface of estuarine mudflats or very shallow pools.

259 Calcium signalling in response to changes in temperature

P. tricornutum cells expressing the R-GECO1 Ca^{2+} biosensor were perfused with seawater at 260 high or low target temperatures (30 °C or 12 °C). Note that actual temperatures in the 261 perfusion dish differed by ± 2 °C from these target temperatures due to equilibration of the 262 small volume of warm or cold perfusate with room temperature. Actual dish temperatures 263 were therefore recorded and are displayed for all experiments. We routinely observed a single 264 transient [Ca²⁺]_{cyt} elevation in cells exposed to a cold shock from 30 °C to 12 °C (97 % cells, 265 n=63) (Fig 2A). In contrast, cells exposed to a rapid rise in temperature from 12 °C to 30 °C 266 did not show [Ca²⁺]_{cvt} elevations (Fig. 2A). No [Ca²⁺]_{cvt} elevations were observed in cells 267 268 perfused with these solutions after they had been equilibrated to room temperature, indicating 269 that the act of switching between the perfusion solutions does not contribute to the signalling responses (Fig. 2A). Analysis of the spatial characteristics of cold-shock induced $[Ca^{2+}]_{cvt}$ 270 271 elevations indicated that many initiate at the apex of the cell and propagate to the towards the 272 central region (Fig 2B), in a manner similar to those induced by mild hypo-osmotic shock 273 (Helliwell et al., 2021). This suggests that the apices of the cell may play an important role in sensing the temperature changes. Cells exposed to a second cold shock two minutes after a 274 previous cold shock demonstrated $[Ca^{2+}]_{cvt}$ elevations with no significant attenuation in 275 amplitude, although the percentage of cells responding was slightly lower (97% to 81 % of 276 277 cells, n=63) (Supplementary Fig S1).

The $[Ca^{2+}]_{cyt}$ elevations observed during cold shock were represented by a >10-fold increase 278 279 in R-GECO1 fluorescence. Assuming a K_d of 480 nM for R-GECO1 and comparison with published maximum F/F₀ values (Zhao et al., 2011), we estimate that $[Ca^{2+}]_{cyt}$ elevations 280 281 reach concentrations in the µM range, which are likely to be physiologically significant. In addition to these large increases in fluorescence that are attributed to [Ca²⁺]_{cyt} elevations, 282 much smaller changes in the baseline fluorescence of each cell could be observed following 283 changes in temperature (increasing with low temperature and decreasing with high 284 temperature, Supplemental Fig S1). These minor changes most likely represent temperature-285 dependent changes in R-GECO1 fluorescence emission (Ohkura et al., 2012) rather than 286 actual changes in resting Ca^{2+} concentration. Therefore, only the substantial transient 287 increases in fluorescence (F/F₀>1.5) representing large [Ca²⁺]_{cyt} elevations were analysed 288 further. 289

290 $[Ca^{2+}]_{cyt}$ elevations were also observed when a cold shock (30°C to 12 °C) was applied to 291 cells held at 22 °C, rather than 30 °C, indicating that the cold shock response was not a 292 consequence of prior warming of the cells (Supplemental Fig S2). The percentage of cells 293 responding to cold shock was lower in cells held at 22 °C compared to cells held at 30 °C, 294 although this may also be influenced by the lower maximum rate of cooling at 22 °C.

295

296 Rapid cooling is required to elicit a $[Ca^{2+}]_{cyt}$ elevation

We therefore examined the nature of the temperature change required to elicit $[Ca^{2+}]_{cvt}$ 297 298 elevations, by manipulating the flow rate of the perfusion to vary the rate of cooling. Rapid cooling (2.5 °C s⁻¹) resulted in $[Ca^{2+}]_{cyt}$ elevations in 100% of cells examined (n=45), whereas 299 only 7% of cells exhibited a [Ca²⁺]_{cyt} elevation at a cooling rate of 0.4 °C s⁻¹ (n=45) (Fig. 3A-300 B). The amplitude of the $[Ca^{2+}]_{cvt}$ elevations in responding cells closely correlated with the 301 cooling rate, with much larger $[Ca^{2+}]_{cyt}$ elevations observed at rapid cooling rates (Fig. 3C). 302 Examination of a broader range of cooling rates indicated that a cooling rate greater than 1 °C 303 s⁻¹ was required to elicit $[Ca^{2+}]_{cyt}$ elevations in 50% of the population (Fig. 3D). These data 304 suggest that the cold shock-induced [Ca²⁺]_{cvt} elevations can therefore relay information 305 relating to the nature of the stimulus both in terms of the number of cells responding and the 306 nature of the $[Ca^{2+}]_{cvt}$ elevation itself. 307

308 As very low perfusion rates also resulted in a lower overall decrease in temperature (due to 309 equilibration of the perfusate with room temperature), we next examined the absolute

temperature decrease required to initiate signalling. Cells were perfused at 30 °C for 1 minute 310 311 and then perfused at a constant flow rate with cold ASW (4 °C) for different durations to vary the decrease in temperature whilst maintaining similar rates of cooling. A very brief perfusion 312 (4 s) lowered the temperature by 2.4 \pm 0.6 °C but did not induce [Ca²⁺]_{evt} elevations (Fig 3E-313 F). However, the rate of cooling in this treatment was considerably lower than the other 314 315 treatments, due to buffering of the temperature by the residual volume within the perfusion dish (1 mL). Perfusions of a longer duration (7-26 s) resulted in a consistent cooling rate of 316 2.1-2.4 °C s⁻¹. A temperature decrease of 8.8 ± 0.4 °C induced a [Ca²⁺]_{cvt} elevation in 38.4% 317 of cells (n=126), whereas greater decreases in temperature resulted in $[Ca^{2+}]_{cvt}$ elevations in 318 nearly all cells (Fig. 3E-F). The amplitude and duration of [Ca²⁺]_{evt} elevations increased with 319 the greater duration of the temperature decrease (Fig 3G-H). A cooling duration of 26 s did 320 not increase the amplitude of the $[Ca^{2+}]_{cvt}$ elevation beyond those observed at 9 s, but greatly 321 increased the duration of the [Ca²⁺]_{cvt} elevation (Fig 3G-H). Taken together, our results show 322 323 that the cooling rate and the duration of the cold shock influence the amplitude and duration of the $[Ca^{2+}]_{cvt}$ elevation and the percentage of cells responding. 324

325

The cold shock response is conserved in the centric diatom *T. pseudonana* but displays different characteristics

T. pseudonana is a planktonic centric diatom found in marine, estuarine and freshwater 328 329 environments (Alverson et al., 2011), where it is also likely to be exposed to significant changes in temperature. We found that T. pseudonana cells expressing the R-GECO1 330 biosensor exhibited [Ca²⁺]_{cvt} elevations in response to cold shock, with the amplitude of these 331 elevations also dependent on the rate of temperature decrease (Fig. 4A-B). As in P. 332 tricornutum, a control perfusion using ASW media equilibrated to room temperature did not 333 induce $[Ca^{2+}]_{cyt}$ elevations (Fig. 4C). The cold induced $[Ca^{2+}]_{cyt}$ elevations in *T. pseudonana* 334 335 were of a lower amplitude but longer duration than those observed in *P. tricornutum*. The percentage of T. pseudonana cells responding to cold shock was also considerably lower than 336 P. tricornutum (19 % vs. 81 %, respectively), although variable levels of expression of R-337 GECO1 in T. pseudonana likely prevented detection of $[Ca^{2+}]_{cyt}$ elevations in all cells within 338 a field of view (Helliwell et al., 2021) (Fig. 4D-F). Taken together, these findings suggest that 339 cold shock-induced [Ca²⁺]_{cvt} elevations are exhibited by both pennate and centric diatom 340 lineages and may therefore represent a conserved mechanism in many diatom species. 341

343 Cellular mechanisms underlying the cold shock response

We next examined the cellular mechanisms responsible for cold shock Ca^{2+} signalling in *P*. 344 tricornutum. Removal of external Ca²⁺ by perfusion of PtR1 cells with cold Ca²⁺-free ASW 345 completely abolished the $[Ca^{2+}]_{cyt}$ elevations (Fig. 5A). Restoration of external Ca^{2+} to cooled 346 cells did not induce a $[Ca^{2+}]_{cvt}$ elevation. However, when these cells were subsequently 347 warmed to 30 °C and then cooled, $[Ca^{2+}]_{evt}$ elevations were observed in the majority of cells. 348 Thus, the generation of cold-induced $[Ca^{2+}]_{cvt}$ elevation depends on the presence of external 349 Ca^{2+} , and the $[Ca^{2+}]_{cyt}$ elevation is triggered by the rapid drop in temperature rather than low 350 absolute temperature itself. 351

352 P. tricornutum lacks cyclic-gated nucleotide channels, which are important for thermal sensing in plants, although it does possess multiple TRP channels (Verret et al., 2010). The 353 temperature-sensitive TRPM8 channel in animal cells is responsible for cold induced $[Ca^{2+}]_{cvt}$ 354 elevations and can be activated directly by the plant secondary metabolite, menthol (Yin et al., 355 2018). Perfusion of PtR1 cells with 1 mM menthol did not elicit $[Ca^{2+}]_{evt}$ elevations, 356 indicating that this ligand is likely specific to the ion channels involved in animal cold 357 signalling (Fig. 5B). In plant and fungal cells, cold shock induced [Ca²⁺]_{cvt} elevations have 358 been studied through the application of dimethyl sulfoxide (DMSO), which is proposed to 359 mimic cold-induced membrane rigidification (Furuya et al., 2014). DMSO elicited [Ca²⁺]_{cvt} 360 elevations in a dose-dependent manner in P. tricornutum, with 8% and 50% of cells 361 exhibiting Ca^{2+} elevation in response to addition of 1 % and 5% DMSO respectively (n = 24, 362 25) (Fig.5C-D). Ruthenium red (RR) is a non-selective Ca^{2+} channel blocker shown to affect 363 numerous TRP channels including the cold sensitive TRPA1 channel (Andrade et al., 2008; 364 Silva et al., 2015; Christensen et al., 2016). RR also inhibits $[Ca^{2+}]_{cvt}$ elevations in P. 365 tricornutum induced by the resupply of phosphate to phosphate-limited cells, but does not 366 inhibit [Ca²⁺]_{cvt} elevations caused by hypo-osmotic shock (Helliwell et al., 2021; Helliwell et 367 al., 2021). Pre-treatment of PtR1 cells for 5 mins with 5-10 µM RR did not significantly 368 reduce the amplitude of cold-induced Ca²⁺ elevations (Fig 5E-F). However, RR treated cells 369 exhibited a significantly slower response time than non-treated control cells (defined as time 370 371 from stimulus to the initial elevation above the threshold of $F/F_0>1.5$) (Fig. 5G), indicating 372 that whilst RR does not prevent the cold shock response, it may partially inhibit a component of the signalling pathway. 373

374 *P. tricornutum* contains three EukCatA channels, which represent a novel class of voltage-375 gated Na^+/Ca^{2+} channel related to single-domain voltage-gated Na^+ channels in bacteria

(BacNav) (Helliwell et al., 2019). As BacNav channels are temperature sensitive (Arrigoni et 376 377 al., 2016), we examined whether P. tricornutum eukcatA1 knockout mutants exhibited an 378 altered response to cold shock. The percentage of responding cells and the mean maximal amplitude of the $[Ca^{2+}]_{cvt}$ elevations did not differ significantly from control PtR1 cells (Fig. 379 5H-I), indicating that EukCatA1 is not required for cold shock induced Ca²⁺ signalling. 380 Further experiments will be needed to determine whether other candidate ion channels, such 381 as the other EukCatA channels or the diatom TRP channels, contribute to this signalling 382 383 response.

384

385 The cold shock response is not required for growth at low temperatures

We next examined whether the cold shock Ca^{2+} signal was required for *P. tricornutum* cells to survive following a cold shock. We applied cold shocks in the presence and absence of external Ca^{2+} and then monitored their physiology and subsequent growth at 4 °C or 18 °C in the presence of Ca^{2+} (Fig 6A). While cells grew significantly more slowly at 4 °C versus 18 °C, there was no significant impact of inhibiting Ca^{2+} signalling during cold shock on the ability of cold-shocked cells to grow at either 4 °C or 18 °C (Fig 6B).

Growth of *P. tricornutum* at 4 °C promoted the accumulation of the oval morphotype, as reported previously (De Martino et al., 2011) (Fig 6C). Cells acclimated to low temperatures may therefore undergo physiological changes that render them less sensitive to rapid cooling. However, fusiform cells grown at 4 °C for four days still showed a typical cold shock response with no significant difference in the percentage of cells exhibiting a response (Fig. 6D-E).

Taken together, these experiments do not indicate a direct requirement for the $[Ca^{2+}]_{cyt}$ elevations in protection from rapid cooling alone, as inhibition of the signalling response did not adversely affect physiology or growth following a cold shock, and the signalling response was not altered in cells acclimated to low temperatures.

402

403 Interaction between cold and hypo-osmotic shock Ca²⁺ signalling pathways

Diatoms inhabiting inter-tidal regions may regularly experience a cold shock during tidal cycles (Fig 1), but this is unlikely to represent an isolated stressor. In particular, warming of shallow tidal pools can greatly increase their salinity due to evaporation (Firth and Williams, 2009), leading to a significant hypo-osmotic shock when the incoming tide reaches the tidal

408 pool. *P. tricornutum* is highly perceptive to hypo-osmotic shock, exhibiting a large transient 409 $[Ca^{2+}]_{cyt}$ elevation similar to those induced by cold shock (Falciatore et al., 2000; Helliwell et 410 al., 2021). Since cold and hypo-osmotic shocks are likely to regularly co-occur in intertidal 411 environments, we examined cellular Ca^{2+} signalling when these stressors were applied 412 simultaneously.

A relatively mild hypo-osmotic shock (100% ASW to 95% ASW) administered to cells at 25 413 °C resulted in a single $[Ca^{2+}]_{evt}$ elevation, as observed previously (Helliwell et al., 2021) (Fig. 414 7A). When the same hypo-osmotic shock was applied simultaneously with a cold shock (25 415 °C to 10 °C), both the amplitude and duration of the [Ca²⁺]_{cyt} elevations was substantially 416 increased, although the number of cells exhibiting $[Ca^{2+}]_{cvt}$ elevations did not change (Fig. 417 418 7A-C). Hypo-osmotic shocks cause an increase in cell volume in *P. tricornutum*, which likely initiates [Ca²⁺]_{cvt} elevations through the activation of mechanosensitive ion channels 419 (Helliwell et al., 2021). However, cell volume did not increase during cold shock 420 (Supplemental Fig. S3), indicating that the rapid cooling does not simply elicit $[Ca^{2+}]_{cvt}$ 421 elevations by mimicking a hypo-osmotic stimulus. 422

A stronger hypo-osmotic shock (100% ASW to 50 % ASW) resulted in a rapid [Ca²⁺]_{cvt} 423 elevation which initiated directly after the stimulus was applied (Fig. 7D). In comparison, 424 application of a cold shock from 34 °C to 8 °C triggered [Ca²⁺]_{cvt} elevations that rose less 425 rapidly and exhibited a longer delay to their initiation (Fig. 7D). Combining both shocks using 426 perfusion with 50% ASW at 10 °C led to biphasic [Ca²⁺]_{cvt} elevations in 71 % of cells (42 427 cells, three separate experiments) (Fig. 7D). These consisted of a very rapid initial peak in 428 $[Ca^{2+}]_{cvt}$, followed by a second peak around 3 s later, which was of greater amplitude than the 429 first peak in the majority of cells (24 out of 30). The mean maximal amplitude of the $[Ca^{2+}]_{cvt}$ 430 elevations caused by the three different treatments were all significantly different from each 431 other, with the cold shock alone causing the lowest and the combined cold- and hypo-osmotic 432 shock causing the highest $[Ca^{2+}]_{cvt}$ elevations (Fig. 7E). 433

Taken together, $[Ca^{2+}]_{cyt}$ elevations induced by hypo-osmotic shock exhibit significant differences in amplitude and timing in the presence of a simultaneous cold shock. This indicates that the cold shock stimulus is additive and of sufficient magnitude to influence cellular Ca²⁺ signalling during hypo-osmotic stress. We therefore investigated whether Ca²⁺ signalling during cold shock may influence the short -term survival of *P. tricornutum* under hypo-osmotic stress.

440

441 Simultaneous cold shock enhances survival during hypo-osmotic shock

442 Cells were treated with 25% ASW to administer a strong hypo-osmotic shock at control and low temperatures in the presence or absence of external Ca²⁺. Cell viability was determined 443 after 3 min by staining with Sytox-Green. Administration of a cold shock alone, either in the 444 presence or absence of external Ca^{2+} , did not reduce cell viability (Fig 8). Application of a 445 strong hypo-osmotic shock (25% ASW) significantly reduced cell viability, and this effect 446 was greater following the removal of external Ca²⁺, supporting our previous observations that 447 Ca^{2+} signalling is required for osmoregulation and volume control in *P. tricornutum* 448 (Helliwell et al., 2021). Surprisingly, application of 25% ASW in combination with a cold 449 shock (4 °C) led to a substantial reduction in cell mortality caused by hypo-osmotic shock 450 (compared to the control temperature, 21°C). This effect was reduced by inhibiting Ca²⁺ 451 signalling, although cell viability remained higher than at control temperature. Our data 452 453 therefore indicate that rapid cooling has an important beneficial influence on the survival of P. 454 tricornutum cells during a hypo-osmotic shock.

455 Cold shock is associated with Ca²⁺-dependent K⁺ efflux

Ca²⁺-dependent K⁺ efflux plays an essential role in cellular volume control in *P. tricornutum* 456 during hypo-osmotic shock (Helliwell et al., 2021). We therefore tested whether the $[Ca^{2+}]_{cvt}$ 457 elevations induced by cold shock also resulted in a K⁺ efflux that could influence cellular 458 osmolarity. We settled a mono-layer of P. tricornutum cells onto a microscopy dish and used 459 a K⁺-selective microelectrode to measure changes in extracellular K⁺ in the immediate 460 vicinity of these cells. The cells were perfused with ASW at 25 °C, before rapidly switching 461 to 12 °C. In each case, the cold shock induced a clear increase in extracellular K⁺ around the 462 P. tricornutum cells (Fig 9A-C). Application of a cold shock in the absence of external Ca²⁺ 463 greatly reduced K^+ efflux from the cells, indicating that the K^+ efflux is Ca²⁺-dependent. Very 464 little change in extracellular K⁺ was observed during a cold shock in the absence of cells, 465 indicating that the performance of the K⁺-selective microelectrode was not affected by the 466 change in temperature. We conclude that cold shock induces Ca^{2+} -dependent K⁺ efflux in P. 467 tricornutum cells, which may contribute to volume regulation during a simultaneous hypo-468 469 osmotic shock.

472 Discussion

This study shows that physiologically significant transient $[Ca^{2+}]_{cvt}$ elevations are a consistent 473 response to the rapid cold shocks likely to be experienced by intertidal diatoms. By using a 474 continuous perfusion system, our study was able to avoid a shear-related [Ca²⁺]_{cvt} response, 475 which may have masked a cold $[Ca^{2+}]_{cvt}$ response in earlier investigations using P. 476 tricornutum expressing acquorin (Falciatore et al., 2000). The cold-induced [Ca²⁺]_{cvt} 477 478 elevations are shown to be specifically involved in sensing the rate of cooling rather than the absolute temperature. A similar dependence of the amplitude of $[Ca^{2+}]_{cvt}$ elevations on the rate 479 of cooling has been observed in Arabidopsis, which showed [Ca²⁺]_{cvt} elevations at cooling 480 rates down to 0.05 °C s⁻¹, (Plieth et al., 1999) indicating greater sensitivity of Arabidopsis to 481 slower cooling rates. P. tricornutum and T. pseudonana did not show a Ca^{2+} signalling 482 response to rapid warming, suggesting that the Ca²⁺ signalling pathways of animals and plants 483 in response to elevated temperatures are not conserved in diatoms. Diatoms therefore likely 484 485 use alternative cellular mechanisms for thermosensation in response to rapid heat shock, although as only short-term temperature increases were evaluated in our study, we cannot rule 486 out a potential role for Ca^{2+} signalling in response to longer term temperature increases. 487

Cold shock-induced $[Ca^{2+}]_{cyt}$ elevations in *P. tricornutum* do not play an obvious role in 488 acclimation to low temperatures. We found no longer-term growth effects of experimentally 489 blocking the cold shock Ca²⁺ signal. Cold signalling in *P. tricornutum* therefore differs from 490 plants and insects (Knight and Knight, 2012; Teets et al., 2013), in which the [Ca²⁺]_{cvt} 491 elevations play a direct role in acclimation to lower temperatures. The $[Ca^{2+}]_{evt}$ response in P. 492 493 tricornutum is specifically induced by rapid cooling, which points to a potential role in short-494 term regulation of cellular processes rather than longer term acclimation to a change in 495 temperature. Of particular interest is the interaction between cold shock and osmotic shock, 496 since inter-tidal organisms are often likely to experience these stresses simultaneously, during 497 an incoming tide or rain precipitation (Lewin and Guillard, 1963; Kirst, 1990). Given the nature of the osmotic and cold shock Ca^{2+} signals identified in *P. tricornutum*, it is most likely 498 that they involve distinct sensory pathways, as evidenced by their additive nature and the 499 appearance of a biphasic $[Ca^{2+}]_{cvt}$ elevation when cells were treated with simultaneous cold 500 and osmotic shocks. Whether these distinct responses represent Ca²⁺ entry through different 501 Ca^{2+} channels or are due to sequential activation of the same Ca^{2+} channel by different stimuli 502 with little or no refractory period remains to be determined, although it is worthy of note that 503 both the osmotic (Helliwell et al., 2021) and the cold-induced Ca²⁺ signals initiate at the cell 504

apices (Fig 2B). Cold and osmotic Ca^{2+} signals also both require the presence of external Ca^{2+} , indicating a shared requirement for plasma membrane Ca^{2+} channels, at least in the initiation of the $[Ca^{2+}]_{cyt}$ elevation.

The protective effect of cold shock on survival of P. tricornutum in response to severe 508 osmotic shock may arise directly from cooperative Ca²⁺ signalling (Supplemental Fig. S4). 509 The hypo-osmotic shock induced $[Ca^{2+}]_{cyt}$ elevations lead to rapid efflux of K⁺ in P. 510 tricornutum, which restricts cell volume increase and prevents bursting (Helliwell et al., 511 2021). The results here strongly suggest that cold-induced $[Ca^{2+}]_{cvt}$ elevations may also act 512 directly to trigger K^+ efflux from the cytosol, for example through the activation of Ca^{2+} -513 dependent K⁺ channels. Whether the rapid loss of K⁺ plays a physiological role in acclimation 514 515 to low temperature is unclear, but it would clearly serve to lower the osmolarity of the cell. Given the frequent co-occurrence of cold and hypo-osmotic shocks, the cold-induced $[Ca^{2+}]_{cvt}$ 516 elevations may therefore function primarily to support osmoregulation. Rapid cooling does 517 not appear to adversely harm the cell when Ca^{2+} signalling is inhibited, whereas a severe 518 hypo-osmotic shock will lead to cell bursting within seconds if cell volume is not controlled 519 520 (Helliwell et al., 2021).

521 Osmoregulation in response to hypo-osmotic stress in diatoms (and most other eukaryotes) is most likely initiated by activation of mechanosensitive channels due to the increase in cell 522 volume (Helliwell et al., 2021). Mechanosensitive channels only activate when the membrane 523 is under tension, i.e. when swelling has already occurred, and cell viability is therefore under 524 immediate threat if rapid osmoregulation cannot be achieved. The K⁺ efflux in response to a 525 cold shock would allow the cell to reduce its osmolarity even if this critical increase in 526 membrane tension is not perceived. By associating K^+ efflux with an additional stimulus that 527 528 commonly co-occurs with hypo-osmotic shock, diatoms can augment the osmoregulatory response and help prevent cell swelling to critical levels. Consistent with this hypothesis, 529 cold-induced [Ca²⁺]_{cvt} elevations were only associated with very rapid cooling. A more 530 gradual exposure to hypo-osmotic stress conveys a much lower risk of cell bursting, reducing 531 532 the need to augment the osmoregulatory response.

We should also consider that low temperature may have a direct effect on reducing mortality during hypo-osmotic stress that is independent of the signalling component, for example by increasing cell wall rigidity. However, the protective effect of cold shock in the absence of Ca^{2+} was small compared to the much greater reduction in mortality in the presence of external Ca^{2+} . We were unable to identify pharmacological inhibitors to selectively inhibit

either osmotic or cold associated Ca²⁺ signalling and the removal of external Ca²⁺ completely
inhibited both signalling pathways. Dissecting the individual contributions of these signalling
pathways to cell survival during simultaneous shocks is therefore not currently easily
achieved. Selective inactivation of the underlying molecular mechanisms through genetic
approaches will most likely be required to fully understand the nature of the cross-talk
between the signalling pathways.

544 Cellular responses to stressors are commonly examined in isolation in the laboratory in order 545 to simplify the elucidation of the signalling pathways responsible. However, organisms often 546 have to respond to inputs from multiple stimuli simultaneously in their natural environment, 547 leading to cross-talk between signalling pathways. Cross-talk in cell signalling can occur 548 when two distinct stimuli trigger a shared cellular response that confers tolerance to both 549 stressors. This may involve activation of a common receptor or activation of independent 550 receptors that converge on a specific node in the signalling pathway (Knight and Knight, 551 2001). Cross-talk with temperature sensing is likely to have evolved when another stress 552 occurs simultaneously with temperature or with a predictable temporal link (i.e. one stimulus 553 consistently precedes the other) (Sinclair et al., 2013). In the case of the inter-tidal zone, many environmental parameters will exhibit a degree of co-variance associated with tidal immersion 554 555 and emersion. It seems likely that organisms inhabiting these environments have developed 556 mechanisms of cross-talk in their pathways of environmental perception that enable them optimise their physiological responses. 557

558 There are multiple examples of cross-talk between temperature and osmotic stress signalling 559 pathways in other eukaryotes. In plants, freezing temperatures can lead to cellular water loss 560 due to external ice formation and many of the genes within the cold-inducible COR regulon 561 are also inducible by drought (Boyce et al., 2003). The cold-responsive CBF/DREB1 and 562 drought-responsive DREB2 transcription factors both bind to a common promoter element 563 (DRE), leading to convergence of the cold and drought signalling pathways (Boyce et al., 2003). Overexpression of the cold-responsive DREB1A transcription factor in Arabidopsis 564 565 resulted in enhanced tolerance to both freezing and drought stress (Liu et al., 1998). In 566 addition, Arabidopsis plants treated with the phytohormone abscisic acid, which plays a 567 primary role in drought tolerance, also show enhanced freezing tolerance (Mantyla et al., 1995). Cross-talk between temperature and osmotic stress signalling pathways have also been 568 569 documented in yeast. Saccharomyces cerevisiae exhibits a high osmolarity (HOG) response to 570 hyper-osmotic stress that results in increased production of the compatible solute, glycerol.

The HOG response is mediated by a mitogen-activated protein kinase (MAPK) pathway that can also be activated by other stimuli including both cold and heat shocks. Heat shock activates the HOG pathway indirectly by stimulating loss of glycerol, leading to hyperosmotic stress (Winkler et al., 2002; Dunayevich et al., 2018).

Our results indicate that cross-talk between Ca²⁺-mediated cellular signalling mechanisms is 575 576 an important consideration in the response of marine organisms to multiple stressors. Whilst our results are discussed primarily in the context of the inter-tidal zone where rapid 577 578 substantial changes in temperature are a regular occurrence, the conserved nature of coldinduced Ca^{2+} signalling in *T. pseudonana* suggests that this pathway may be important more 579 widely in diatom ecology. The cold-induced $[Ca^{2+}]_{cvt}$ elevations in *T. pseudonana* exhibit 580 581 different characteristics from *P. tricornutum* that may reflect differences in their physiological 582 response. Planktonic diatoms will undoubtedly encounter significant fluctuations in temperature and salinity in near-shore and estuarine environments or when they are mixed 583 584 through the thermocline, although the magnitude and rate of the temperature changes are likely to be lower. Diatoms inhabiting sea ice environments may also experience rapid 585 586 changes in temperature and salinity e.g. during flushing of hyper-saline brine channels with melt water (Mock and Junge, 2007). Future elucidation of the mechanisms of cross talk in 587 588 these signalling pathways will allow us to understand how diatoms successfully integrate inputs from multiple environmental stimuli, which has likely played a major role in their 589 590 success in diverse and highly dynamic environmental regimes.

591 Acknowledgements:

592 The work was supported by an ERC Advanced Grant to CB (ERC-ADG-670390) and a 593 NERC award to GLW (NE/V013343/1).

594

595 Author contributions

596 CB, FK and GLW conceived the study. FK performed the majority of the experimental 597 analyses, including all imaging experiments. KEH contributed to measurements of osmotic 598 stress. AC performed K^+ microelectrode measurements. HPW and NM performed the 599 environmental monitoring. TM, AH and TG contributed to the transformation of *T*. 600 *pseudonana* with R-GECO1. GLW, CB and FK analysed the data and wrote the manuscript.

602 Figures

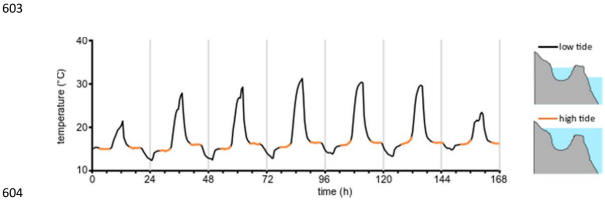


Figure 1: Temperature fluctuations in the inter-tidal zone. An example of temperature fluctuations measured in a temperate coastal rock pool (Looe, Cornwall, UK) over the course of seven days in summer (01/07/2019-07/07/2019). Orange colour indicates periods at which the pool was immersed by the high tide (approx. duration of immersion 5 h). Significant excursions from the sea temperature occur when the rock pool is isolated from the bulk seawater at low tide (black traces). Rapid cooling (30 °C to 15 °C) occurs when the incoming tide reaches the pool.

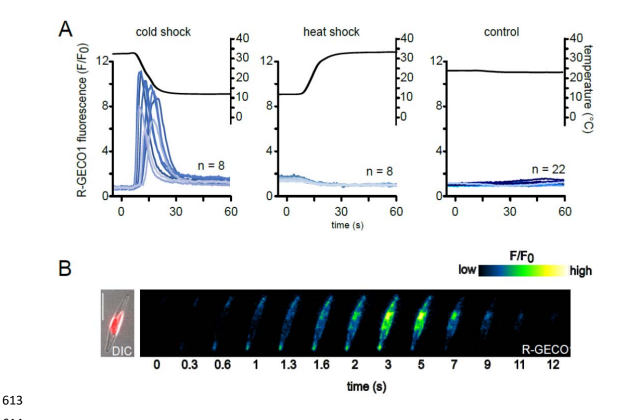


Figure 2: *P. tricornutum* exhibits cytosolic Ca²⁺ ([Ca²⁺]_{cyt}) elevations in response to rapid 615 cooling. A) Eight representative fluorescence ratio traces (F/F₀, blue lines) of *P. tricornutum* 616 cells expressing R-GECO1 representing changes in cytosolic Ca²⁺. Cells were perfused with 617 ASW of different temperatures to cause rapid temperature shifts (black line). Cold shock 30 618 °C to 12 °C, heat shock 12 °C to 30 °C or control 22 °C to 22 °C. B) False colour images of a 619 PtR1 cell exhibiting a $[Ca^{2+}]_{cvt}$ elevation in response to cold shock. The temperature decrease 620 begins at t = 0 s. Note that the $[Ca^{2+}]_{cvt}$ elevations initiate at the tips of the cell and spread 621 622 towards the central region. Left panel indicates a DIC image overlaid with chlorophyll 623 autofluorescence. Bar represents 10 µm.

bioRxiv preprint doi: https://doi.org/10.1101/2022.03.11.483981; this version posted March 12, 2022. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

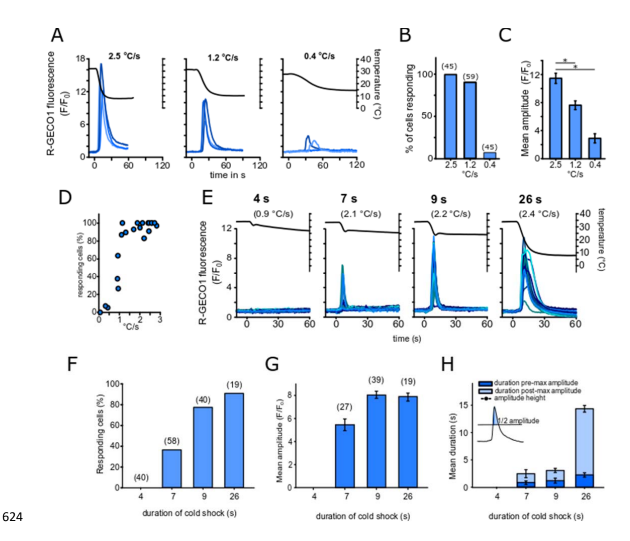


Figure 3: The cold shock Ca^{2+} response depends on the rate of change of temperature. 625 A) R-GECO1 fluorescence in P. tricornutum in response to cold shock administered at 626 627 different cooling rates. As cooling rates were non-linear the maximal cooling rate for each 628 treatment was calculated for comparisons. Three representative traces are shown. B) The percentage of cells exhibiting a $[Ca^{2+}]_{cvt}$ elevation (F/F₀>1.5) at different cooling rates. Total 629 number of cells examined are shown in parentheses, from a minimum of two separate 630 experimental treatments. C) Mean maximal amplitude of $[Ca^{2+}]_{cvt}$ elevations from responsive 631 cells in (B). * indicates a significant difference (One way ANOVA on Ranks P>0.001, 632 Dunn's post-hoc test P>0.001). n=31,30 and 3 for 2.5, 1.2 and 0.4 °C s⁻¹ respectively. Error 633 bars = SE. D) Percentage of cells responding to cold shock with a $[Ca^{2+}]_{cvt}$ elevation across a 634 635 broader range of maximum cooling rates. The data represent 21 independent experiments, with a mean of 38 cells examined for each data point (minimum 12, maximum 123 cells). E) 636 $[Ca^{2+}]_{cvt}$ elevations in response to different durations of cooling applied with a constant flow 637 rate (16 mL min⁻¹). 20 representative traces from PtR1 cells are shown, with greater $[Ca^{2+}]_{cvt}$ 638 639 elevations observed under increasing durations of cold shock. The maximum rate of

temperature decrease ($\Delta T s^{-1}$) is shown in parantheses. Data for 4, 7, 9 and 26 s of cold shock 640 641 duration were compiled from 2, 3, 2 and 1 individual experiments, respectively. F) The percentage of cells exhibiting a [Ca²⁺]_{cyt} elevation in response to cold shock for the 642 experiment described in (E).G) Mean maximal amplitude of $[Ca^{2+}]_{cyt}$ elevations in response to 643 cold shock for the responding cells shown in (F). H) Duration of $[Ca^{2+}]_{cyt}$ elevations (shown 644 as full width at half maximum amplitude) in relation of the duration of cold stimulus. The 645 duration of $[Ca^{2+}]_{cvt}$ elevations is greatest at the 26 s cold shock. The duration is divided into a 646 pre- and post-maximal amplitude component to show that the post-maximal amplitude (tail) 647 components of the $[Ca^{2+}]_{cyt}$ elevation is greatly extended under the 26 s cold shock. 648

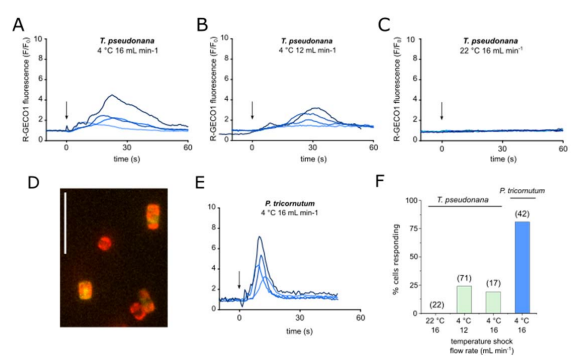


Figure 4: Thalassiosira pseudonana also shows cold-induced [Ca²⁺]_{cyt} elevations. A) 650 Fluorescence ratio of T. pseudonana cells expressing cytosolic R-GECO1 in response to a 651 cold shock (from 30 to 10 °C). For these experiments the temperature in the dish was not 652 653 monitored, so perfusion flow rate is shown to indicate rate of cold shock. Arrow indicates onset of cold stimulus. Four representative traces are shown. B) As in (A) but at a slower flow 654 rate. C) Treatment control using perfusion of ASW at room temperature. D) Fluorescence 655 656 image of T. pseudonana cells expressing R-GECO1 (yellow) overlaid with chlorophyll 657 autofluorescence (red). Scale bar represents 20 µm. E) P. tricornutum cold shock response under identical treatment as in A for comparison. F) Percentage of cells exhibiting $[Ca^{2+}]_{cvt}$ 658 elevations. Values in parentheses denote n. 659

660



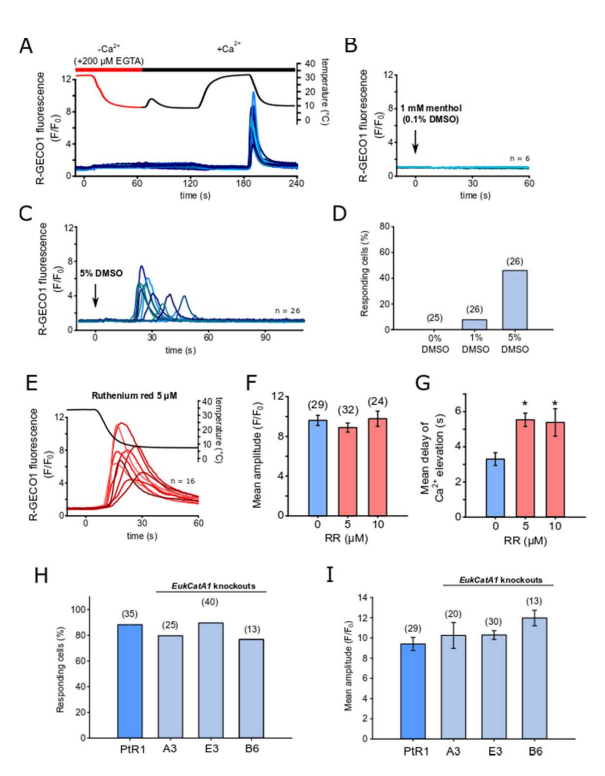
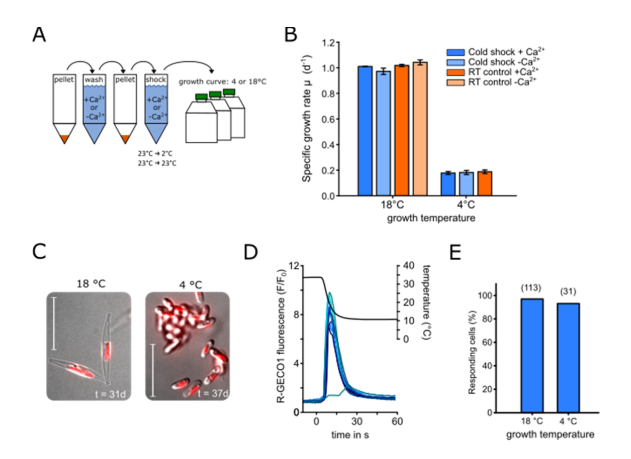




Figure 5: Cellular mechanisms of cold shock induced $[Ca^{2+}]_{cyt}$ elevations. A) R-GECO1 fluorescence ratio (F/F₀) from a cold shock applied to PtR1 cells using ASW without Ca²⁺ + 200 μ M EGTA (Methods). No $[Ca^{2+}]_{cyt}$ elevations can be observed during the cold shock or when Ca²⁺ was restored to cold cells (perfused with cold ASW with Ca²⁺). However, $[Ca^{2+}]_{cyt}$ elevations were observed during a subsequent cold shock applied with standard ASW (i.e.

with 10 mM CaCl₂). Note the minor temperature increase at 70 s is due to a slight warming of 668 cold ASW+Ca²⁺ media in the perfusion system. 23 representative traces are shown, three 669 additional experiments were performed with identical results. B) PtR1 fluorescence in 670 671 response to ASW containing 1 mM menthol (including 0.1% DMSO as solvent carrier). Six representative traces are shown. C) R-GECO1 fluorescence ratio of PtR1 cells perfused with 672 ASW + 5% DMSO. **D**) Percentage of cells exhibiting $[Ca^{2+}]_{evt}$ elevations in response to 673 DMSO. E) The effect of cold shock on PtR1 cells pre-treated with the Ca^{2+} channel blocker 674 ruthenium red (5 µM final, 5 min pre-incubation). 16 representative traces are shown. F) 675 Mean amplitude (\pm SE) of responding cells treated with ruthenium red (5 μ M or 10 μ M) 676 compared to untreated control cells. No significant differences were observed (one-way 677 ANOVA). G) Mean timing (\pm SE) of the maximal amplitude of $[Ca^{2+}]_{cvt}$ elevations for cells 678 treated with ruthenium red (P = <0.01, one-way ANOVA, Holm-Sidak post-hoc test). H) 679 Percentage of cells showing $[Ca^{2+}]_{cvt}$ elevations in response to cold shock in control and three 680 681 independent eukcatal mutant strains (30 to 10 °C). Data represent a minimum of two 682 independent experiments per strain (one-way ANOVA). I) Mean maximal amplitude (±SE) of [Ca²⁺]_{cvt} elevations in *eukcatA1* mutants in response to cold shock. No significant differences 683 684 were observed (one-way ANOVA on Ranks).



686

Figure 6: The role of the cold shock response in cold tolerance. A) Schematic diagram 687 showing the workflow for an experiment examining the impact of Ca^{2+} -dependent cold shock 688 signalling on P. tricornutum cold tolerance. Cells were harvested and washed in ASW 689 containing 10 mM Ca²⁺ or no Ca²⁺ (ASW -Ca²⁺ +200 µM EGTA). Cells were pelleted once 690 again and exposed to cold shock with or without Ca^{2+} . Cells were then grown at control (18) 691 °C) and cold (4 °C) conditions in standard ASW (i.e. with 10 mM CaCl₂) to examine cold 692 tolerance. B) Growth rate of P. tricornutum cultures after cold shock treatment. Mean specific 693 growth rates were calculated from day 0-5 and 12-30 for 18 °C and 4 °C, respectively. Note 694 that for growth at 4 °C, a no shock (RT) control in the absence of Ca²⁺ was not included. No 695 significant differences were observed between treatments at each temperature (one-way 696 697 ANOVA). Error bars represent SE. C) DIC images of PtR1 cells grown at 18 or 4 °C. Oval cells predominate in cells grown at 4 C for extended periods. Red = chlorophyll auto-698 fluorescence, bar = 20 μ m. D) Cold-acclimated PtR1 cells still exhibit a response to cold 699 shock. Representative R-GECO1 fluorescence ratio traces from PtR1 cells grown at 4 °C for 4 700 701 days. Cells were briefly warmed to 30 °C before a cold shock was applied. E) The percentage 702 of cells responding to cold shock as function of acclimation temperature. The results were

- generated from four separate experiments with maximum temperature drop-rates between 2.2
- **704** $3.2 \, \text{C}^{\circ} \, \text{s}^{-1}$.

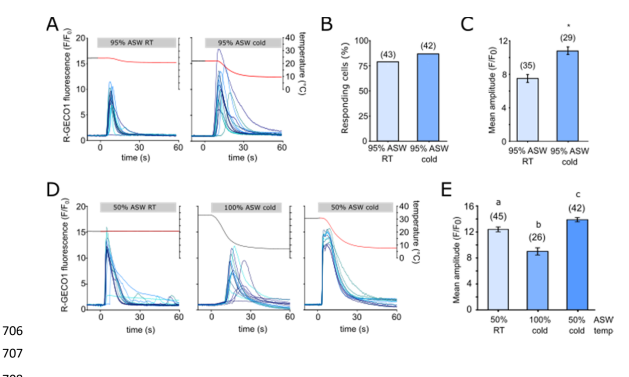
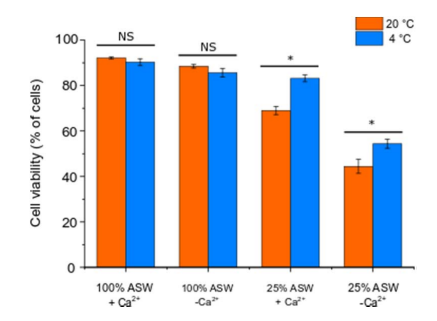


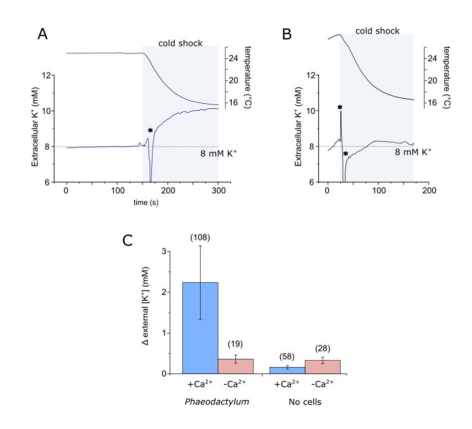


Figure 7: Interactions between the cold shock and hypo-osmotic shock Ca²⁺ signalling 709 pathways. A) R-GECO1 fluorescence ratio (F/F_0) of PtR1 cells in response to a mild hypo-710 711 osmotic shock (95% ASW, left panel) or a simultaneous hypo-osmotic and cold shock (10 °C decrease, right panel). 12 representative traces are shown. B) Percentage of cells exhibiting 712 $[Ca^{2+}]_{cvt}$ elevations for the experiment described in (A). Data are compiled from a minimum 713 of two independent treatments. C) Mean amplitude of $[Ca^{2+}]_{cvt}$ elevations from responding 714 cells in (B). The two treatments are significantly different (Student's t-test P < 0.001). D) R-715 GECO1 fluorescence ratio of PtR1 cells in response to stronger simultaneous cold- and hypo-716 osmotic shocks. Cells were treated with a single hypo-osmotic shock (50% ASW), a single 717 cold shock (10 °C) or a simultaneous cold- and hypo-osmotic shock (50% ASW, 10 °C). 13 718 representative traces are shown. E) Mean maximal amplitude of cells exhibiting $[Ca^{2+}]_{cvt}$ 719 elevations in (D). For biphasic peaks the higher amplitude was chosen. The data represent the 720 combination of at least three independent experiments per treatment. Letters represent 721 significant differences between treatments (1-way Kruskal-Wallis ANOVA Ranks P < 0.001, 722 with Dunn post hoc). 723



724

Figure 8: Simultaneous cold shock reduces mortality associated with hypo-osmotic 725 726 shock. Cell viability (measured by exclusion of Sytox Green stain) was determined in P. 727 tricornutum cells 3 min after exposure to a severe hypo-osmotic shock (25% ASW) with or without simultaneous cold shock (4 °C ASW). The presence or absence of external Ca²⁺ was 728 used to establish the effect of inhibiting Ca^{2+} signalling during the applied shocks. The 729 decrease in cell viability due to a hypo-osmotic shock (25% ASW) is significantly reduced 730 when a simultaneous cold shock is applied. Three replicates were performed for each 731 treatment, with at least 100 cells counted for each replicate. Significant differences due to 732 temperature are marked with * (P<0.05 1-way ANOVA with Holm-Sidak post-hoc test). The 733 experiment was repeated four times with similar results each time. Error bars represent SE. 734



735

736

Figure 9: Cold shock induces a Ca^{2+} -dependent K⁺ efflux. A) K⁺ efflux from P. 737 tricornutum cells during a cold shock. A K⁺ microelectrode was placed adjacent to densely-738 packed *P. tricornutum* cells to measure K+ in the immediate vicinity of the cell. A cold shock 739 was applied by perfusion. The increase in extracellular K^+ is the result of K^+ efflux from the 740 cells. The temperature in the dish is also shown (upper trace). **B**) Extracellular K^+ during a 741 cold shock in the absence of external Ca^{2+} (perfusion with ASW- Ca^{2+} + 200 µM EGTA). C) 742 Mean change in extracellular K⁺ around *P. tricornutum* cells during a cold shock. White bars 743 indicate control experiments where the experimental set up was identical, but no P. 744 tricornutum cells were present in order to assess whether the performance of the K⁺ 745 microelectrode was influenced by temperature. The total number of replicates for each 746 treatment are shown in parentheses, error bars = SE. 747

Parsed Citations

Alverson AJ, Beszteri B, Julius ML, Theriot EC (2011) The model marine diatom Thalassiosira pseudonana likely descended from a freshwater ancestor in the genus Cyclotella. BMC Evol Biol 11: 125

Google Scholar: Author Only Title Only Author and Title

Anderson SI, Rynearson TA (2020) Variability approaching the thermal limits can drive diatom community dynamics. Limnology and Oceanography 65: 1961-1973

Google Scholar: Author Only Title Only Author and Title

Andrade EL, Luiz AP, Ferreira J, Calixto JB (2008) Pronociceptive response elicited by TRPA1 receptor activation in mice. Neuroscience 152: 511-520

Google Scholar: <u>Author Only Title Only Author and Title</u>

Arrigoni C, Rohaim A, Shaya D, Findeisen F, Stein RA, Nurva SR, Mishra S, McHaourab HS, Minor DL, Jr. (2016) Unfolding of a temperature-sensitive domain controls voltage-gated channel activation. Cell 164: 922-936 Google Scholar: Author Only Title Only Author and Title

Baker KG, Geider RJ (2021) Phytoplankton mortality in a changing thermal seascape. Glob Chang Biol Google Scholar: <u>Author Only Title Only Author and Title</u>

Boyce JM, Knight H, Deyholos M, Openshaw MR, Galbraith DW, Warren G, Knight MR (2003) The sfr6 mutant of Arabidopsis is defective in transcriptional activation via CBF/DREB1 and DREB2 and shows sensitivity to osmotic stress. Plant J 34: 395-406 Google Scholar: Author Only Title Only Author and Title

Christensen AP, Akyuz N, Corey DP (2016) The outer pore and selectivity filter of TRPA1. PLoS One 11: e0166167 Google Scholar: <u>Author Only Title Only Author and Title</u>

Clapham DE, Miller C (2011) A thermodynamic framework for understanding temperature sensing by transient receptor potential (TRP) channels. Proc Natl Acad Sci U S A 108: 19492-19497 Google Scholar: Author Only Title Only Author and Title

Cui Y, Lu S, Li Z, Cheng J, Hu P, Zhu T, Wang X, Jin M, Wang X, Li L, Huang S, Zou B, Hua J (2020) Cyclic nucleotide-gated ion channels 14 and 16 promote tolerance to heat and chilling in rice. Plant Physiol 183: 1794-1808 Google Scholar: Author Only Title Only Author and Title

De Martino A, Bartual A, Willis A, Meichenin A, Villazan B, Maheswari U, Bowler C (2011) Physiological and molecular evidence that environmental changes elicit morphological interconversion in the model diatom Phaeodactylum tricornutum. Protist 162: 462-481

Google Scholar: Author Only Title Only Author and Title

De Martino A, Meichenin A, Shi J, Pan KH, Bowler C (2007) Genetic and phenotypic characterization of Phaeodactylum tricornutum (Bacillariophyceae) accessions. Journal of Phycology 43: 992-1009 Google Scholar: Author Only Title Only Author and Title

Dunayevich P, Baltanas R, Clemente JA, Couto A, Sapochnik D, Vasen G, Colman-Lerner A (2018) Heat-stress triggers MAPK crosstalk to turn on the hyperosmotic response pathway. Sci Rep 8: 15168

Google Scholar: <u>Author Only Title Only Author and Title</u>

Falciatore A, d'Alcala MR, Croot P, Bowler C (2000) Perception of environmental signal by a marine diatom. Science 288: 2363-2366

Google Scholar: Author Only Title Only Author and Title

Firth LB, Williams GA (2009) The influence of multiple environmental stressors on the limpet Cellana toreuma during the summer monsoon season in Hong Kong. Journal of Experimental Marine Biology and Ecology 375: 70-75

Google Scholar: Author Only Title Only Author and Title

Furuya T, Matsuoka D, Nanmori T (2014) Membrane rigidification functions upstream of the MEKK1-MKK2-MPK4 cascade during cold acclimation in Arabidopsis thaliana. FEBS Lett 588: 2025-2030

Google Scholar: Author Only Title Only Author and Title

Gattuso JP, Magnan A, Bille R, Cheung WWL, Howes EL, Joos F, Allemand D, Bopp L, Cooley SR, Eakin CM, Hoegh-Guldberg O, Kelly RP, Portner HO, Rogers AD, Baxter JM, Laffoley D, Osborn D, Rankovic A, Rochette J, Sumaila UR, Treyer S, Turley C (2015) Contrasting futures for ocean and society from different anthropogenic CO2 emissions scenarios. Science 349 Google Scholar: <u>Author Only Title Only Author and Title</u>

Gruber N, Boyd PW, Frolicher TL, Vogt M (2021) Biogeochemical extremes and compound events in the ocean. Nature 600: 395-407

Google Scholar: Author Only Title Only Author and Title

bioRxiv preprint doi: https://doi.org/10.1101/2022.03.11.483981; this version posted March 12, 2022. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license. Guillard RL (1975) Culture of phytoplankton for feeding marine invertebrates. Culture of Marine Invertebrate Animals: 29-60

Google Scholar: Author Only Title Only Author and Title

Harley CDG, Hughes AR, Hultgren KM, Miner BG, Sorte CJB, Thornber CS, Rodriguez LF, Tomanek L, Williams SL (2006) The impacts of climate change in coastal marine systems. Ecology Letters 9: 228-241 Google Scholar: Author Only Title Only Author and Title

Helliwell KE, Chrachri A, Koester JA, Wharam S, Verret F, Taylor AR, Wheeler GL, Brownlee C (2019) Alternative mechanisms for fast Na+/Ca2+ signaling in eukaryotes via a novel class of single-domain voltage-gated channels. Curr Biol 29: 1503-1511 e1506 Google Scholar: Author Only Title Only Author and Title

Helliwell KE, Harrison EL, Christie-Oleza JA, Rees AP, Kleiner FH, Gaikwad T, Downe J, Aguilo-Ferretjans MM, Al-Moosawi L, Brownlee C, Wheeler GL (2021) A Novel Ca2+ signaling pathway coordinates environmental phosphorus sensing and nitrogen metabolism in marine diatoms. Curr Biol 31: 978-989 e974

Google Scholar: Author Only Title Only Author and Title

Helliwell KE, Kleiner FH, Hardstaff H, Chrachri A, Gaikwad T, Salmon D, Smirnoff N, Wheeler GL, Brownlee C (2021) Spatiotemporal patterns of intracellular Ca2+ signalling govern hypo-osmotic stress resilience in marine diatoms. New Phytol 230: 155-170

Google Scholar: Author Only Title Only Author and Title

J C Lewin a, Guillard RRL (1963) Diatoms. Annual Review of Microbiology 17: 373-414 Google Scholar: Author Only Title Only Author and Title

Javaheri N, Dries R, Burson A Stal LJ, Sloot PMA Kaandorp JA (2015) Temperature affects the silicate morphology in a diatom. Scientific Reports 5: 11652.

Google Scholar: Author Only Title Only Author and Title

Kirst GO (1990) Salinity tolerance of eukaryotic marine-algae. Annual Review of Plant Physiology and Plant Molecular Biology 41: 21-53

Google Scholar: Author Only Title Only Author and Title

Knight H, Knight MR (2001) Abiotic stress signalling pathways: specificity and cross-talk. Trends Plant Sci 6: 262-267 Google Scholar: Author Only Title Only Author and Title

Knight H, Trewavas AJ, Knight MR (1996) Cold calcium signaling in Arabidopsis involves two cellular pools and a change in calcium signature after acclimation. Plant Cell 8: 489-503

Google Scholar: Author Only Title Only Author and Title

Knight MR, Knight H (2012) Low-temperature perception leading to gene expression and cold tolerance in higher plants. New Phytol 195: 737-751

Google Scholar: Author Only Title Only Author and Title

Lenzoni G, Knight MR (2019) Increases in absolute temperature stimulate free calcium concentration elevations in the chloroplast. Plant Cell Physiol 60: 538-548

Google Scholar: Author Only Title Only Author and Title

Lewin JC, Guillard RRL (1963) Diatoms. Annual Review of Microbiology 17: 373-414 Google Scholar: Author Only Title Only Author and Title

Liang Y, Koester JA, Liefer JD, Irwin AJ, Finkel ZV (2019) Molecular mechanisms of temperature acclimation and adaptation in marine diatoms. ISME J 13: 2415-2425

Google Scholar: Author Only Title Only Author and Title

Liu Q. Ding Y. Shi Y. Ma L. Wang Y. Song C. Wilkins KA. Davies JM. Knight H. Knight MR. Gong Z. Guo Y. Yang S (2021) The calcium transporter ANNEXIN1 mediates cold-induced calcium signaling and freezing tolerance in plants. EMBO J 40: e104559 Google Scholar: Author Only Title Only Author and Title

Liu Q, Kasuga M, Sakuma Y, Abe H, Miura S, Yamaguchi-Shinozaki K, Shinozaki K (1998) Two transcription factors, DREB1 and DREB2, with an EREBP/AP2 DNA binding domain separate two cellular signal transduction pathways in drought- and lowtemperature-responsive gene expression, respectively, in Arabidopsis. Plant Cell 10: 1391-1406 Google Scholar: Author Only Title Only Author and Title

Ma Y, Dai X, Xu Y, Luo W, Zheng X, Zeng D, Pan Y, Lin X, Liu H, Zhang D, Xiao J, Guo X, Xu S, Niu Y, Jin J, Zhang H, Xu X, Li L, Wang W, Qian Q, Ge S, Chong K (2015) COLD1 confers chilling tolerance in rice. Cell 160: 1209-1221 Google Scholar: Author Only Title Only Author and Title

Malviya S, Scalco E, Audic S, Vincent F, Veluchamy A, Poulain J, Wincker P, Iudicone D, de Vargas C, Bittner L, Zingone A, Bowler C (2016) Insights into global diatom distribution and diversity in the world's ocean. Proc Natl Acad Sci U S A 113: E1516-1525 Google Scholar: Author Only Title Only Author and Title

available under aCC-BY-NC-ND 4.0 International license. Mantyla E, Lang V, Palva ET (1995) Role of abscisic acid in drought-induced freezing tolerance, cold acclimation, and accumulation of LT178 and RAB18 proteins in Arabidopsis thaliana. Plant Physiol 107: 141-148

Google Scholar: Author Only Title Only Author and Title

Mock T, Junge K (2007) PSYCHROPHILIC DIATOMS: Mechanisms for survival in freeze-thaw cycles. In J Seckbach, ed, Agae and Cyanobacteria in Extreme Environments. Springer, pp 343-364

Google Scholar: Author Only Title Only Author and Title

Montagnes DJS, Franklin DJ (2001) Effect of temperature on diatom volume, growth rate, and carbon and nitrogen content: Reconsidering some paradigms. Limnology and Oceanography 46: 2008-2018 Google Scholar: Author Only Title Only Author and Title

Ohkura M, Sasaki T, Sadakari J, Gengyo-Ando K, Kagawa-Nagamura Y, Kobayashi C, Ikegaya Y, Nakai J (2012) Genetically encoded green fluorescent Ca2+ indicators with improved detectability for neuronal Ca2+ signals. PLoS One 7: e51286 Google Scholar: Author Only Title Only Author and Title

Plieth C, Hansen UP, Knight H, Knight MR (1999) Temperature sensing by plants: the primary characteristics of signal perception and calcium response. Plant J 18: 491-497

Google Scholar: Author Only Title Only Author and Title

Pokorna J. Schwarzerova K. Zelenkova S. Petrasek J. Janotova I. Capkova V. Opatrny Z (2004) Sites of actin filament initiation and reorganization in cold-treated tobacco cells. Plant Cell and Environment 27: 641-653 Google Scholar: Author Only Title Only Author and Title

Saidi Y, Finka A Muriset M, Bromberg Z, Weiss YG, Maathuis FJ, Goloubinoff P (2009) The heat shock response in moss plants is regulated by specific calcium-permeable channels in the plasma membrane. Plant Cell 21: 2829-2843 Google Scholar: Author Only Title Only Author and Title

Schaum CE, Buckling A, Smirnoff N, Studholme DJ, Yvon-Durocher G (2018) Environmental fluctuations accelerate molecular evolution of thermal tolerance in a marine diatom. Nat Commun 9: 1719 Google Scholar: Author Only Title Only Author and Title

Sengupta P, Garrity P (2013) Sensing temperature. Curr Biol 23: R304-307 Google Scholar: Author Only Title Only Author and Title

Silva DF, de Almeida MM, Chaves CG, Braz AL, Gomes MA, Pinho-da-Silva L, Pesquero JL, Andrade VA, Leite Mde F, de Abuquerque JG, Araujo IG, Nunes XP, Barbosa-Filho JM, Cruz Jdos S, Correia Nde A, de Medeiros IA (2015) TRPM8 channel activation induced by monoterpenoid rotundifolone underlies mesenteric artery relaxation. PLoS One 10: e0143171 Google Scholar: Author Only Title Only Author and Title

Sinclair BJ, Ferguson LV, Salehipour-shirazi G, MacMillan HA (2013) Cross-tolerance and cross-talk in the cold: relating low temperatures to desiccation and immune stress in insects. Integr Comp Biol 53: 545-556 Google Scholar: Author Only Title Only Author and Title

Smale DA, Wernberg T, Oliver ECJ, Thomsen M, Harvey BP, Straub SC, Burrows MT, Alexander LV, Benthuysen JA, Donat MG, Feng M, Hobday AJ, Holbrook NJ, Perkins-Kirkpatrick SE, Scannell HA, Sen Gupta A, Payne BL, Moore PJ (2019) Marine heatwaves threaten global biodiversity and the provision of ecosystem services. Nature Climate Change 9: 306-+ Google Scholar: Author Only Title Only Author and Title

Souffreau C, Vanormelingen P, Verleyen E, Sabbe K, Vyverman W (2010) Tolerance of benthic diatoms from temperate aquatic and terrestrial habitats to experimental desiccation and temperature stress. Phycologia 49: 309-324 Google Scholar: Author Only Title Only Author and Title

Svensson F, Norberg J, Snoeijs P (2014) Diatom cell size, coloniality and motility: trade-offs between temperature, salinity and nutrient supply with climate change. Plos One 9

Google Scholar: Author Only Title Only Author and Title

Tahtiharju S, Sangwan V, Monroy AF, Dhindsa RS, Borg M (1997) The induction of kin genes in cold-acclimating Arabidopsis thaliana. Evidence of a role for calcium. Planta 203: 442-447 Google Scholar: Author Only Title Only Author and Title

Teets NM, Yi SX, Lee RE, Jr., Denlinger DL (2013) Calcium signaling mediates cold sensing in insect tissues. Proc Natl Acad Sci USA110:9154-9159

Google Scholar: Author Only Title Only Author and Title

Vardi A, Formiggini F, Casotti R, De Martino A, Ribalet F, Miralto A, Bowler C (2006) A stress surveillance system based on calcium and nitric oxide in marine diatoms. Plos Biology 4: 411-419 Google Scholar: Author Only Title Only Author and Title

Verret F, Wheeler G, Taylor AR, Farnham G, Brownlee C (2010) Calcium channels in photosynthetic eukaryotes: implications for

evolution of calcium-based signalling. New Phytol 187: 23-43

Google Scholar: Author Only Title Only Author and Title

Winkler A, Arkind C, Mattison CP, Burkholder A, Knoche K, Ota I (2002) Heat stress activates the yeast high-osmolarity glycerol mitogen-activated protein kinase pathway, and protein tyrosine phosphatases are essential under heat stress. Eukaryot Cell 1: 163-173

Google Scholar: Author Only Title Only Author and Title

Xu H, Ramsey IS, Kotecha SA, Moran MM, Chong JA, Lawson D, Ge P, Lilly J, Silos-Santiago I, Xie Y, DiStefano PS, Curtis R, Clapham DE (2002) TRPV3 is a calcium-permeable temperature-sensitive cation channel. Nature 418: 181-186 Google Scholar: Author Only Title Only Author and Title

Yin Y, Wu M, Zubcevic L, Borschel WF, Lander GC, Lee SY (2018) Structure of the cold- and menthol-sensing ion channel TRPM8. Science 359: 237-241

Google Scholar: Author Only Title Only Author and Title

Zhao Y, Araki S, Wu J, Teramoto T, Chang YF, Nakano M, Abdelfattah AS, Fujiwara M, Ishihara T, Nagai T, Campbell RE (2011) An expanded palette of genetically encoded Ca2+ indicators. Science 333: 1888-1891

Google Scholar: Author Only Title Only Author and Title