



Short-term response of *Emiliana huxleyi* growth and morphology to abrupt salinity stress

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Abstract. The marine coccolithophore species *Emiliana huxleyi* tolerates a broad range of salinity conditions over its near-global distribution, including the relatively stable physiochemical conditions of open-ocean environments and nearshore environments with dynamic and extreme short-term salinity fluctuations. Previous studies show that salinity impacts the physiology and morphology of *E. huxleyi*, suggesting that salinity stress influences the calcification of this globally important species. However, it remains unclear how rapidly *E. huxleyi* responds to salinity changes and therefore whether *E. huxleyi* morphology is sensitive to short-term transient salinity events (such as occur on meteorological timescales) in addition to longer-duration salinity changes. Here, we investigate the real-time growth and calcification response of two *E. huxleyi* strains isolated from shelf sea environments to the abrupt onset of hyposaline and hypersaline conditions over a time period of 156 h (6.5 d). Morphological responses in the size of the cell covering (coccosphere) and the calcium carbonate plates (coccoliths) that form the coccosphere occurred as rapidly as 24–48 h following the abrupt onset of salinity 25 (hyposaline) and salinity 45 (hypersaline) conditions. Generally, cells tended towards smaller coccospheres (−24 %) with smaller coccoliths (−7 % to −11 %) and reduced calcification under hyposaline conditions, whereas cells growing under hypersaline conditions had either relatively stable coccosphere and coccolith sizes (Mediterranean strain RCC1232) or larger coccospheres (+35 %) with larger coccoliths (+13 %) and increased calcification (Norwegian strain PLYB11). This short-term response is consistent with reported coccolith size trends with salinity over longer durations of low- and high-

salinity exposure in culture and under natural-salinity gradients. The coccosphere size response of PLYB11 to salinity stress was greater in magnitude than was observed in RCC1232 but occurred after a longer duration of exposure to the new salinity conditions (96–128 h) compared to RCC1232. In both strains, coccosphere size changes were larger and occurred more rapidly than changes in coccolith size, which tended to occur more gradually over the course of the experiments. Variability in the magnitude and timing of rapid morphological responses to short-term salinity stress between these two strains supports previous suggestions that the response of *E. huxleyi* to salinity stress is strain specific. At the start of the experiments, the light condition was also switched from a light : dark cycle to continuous light, with the aim of desynchronising cell division. As cell density and mean cell size data sampled every 4 h showed regular periodicity under all salinity conditions, the cell division cycle retained its entrainment to pre-experiment light : dark conditions for the entire experiment duration. Extended acclimation periods to continuous light are therefore advisable for *E. huxleyi* to ensure successful desynchronisation of the cell division cycle. When working with phased or synchronised populations, data should be compared between samples taken from the same phase of the cell division cycle to avoid artificially distorting the magnitude or even direction of physiological or biogeochemical response to the environmental stressor.

1 Introduction

Shifts in ocean salinity are an indirect effect of climate change, reflecting changes in the hydrological cycle on seasonal-to-multiannual timescales (precipitation, evaporation, and river runoff) and dynamics in ocean circulation patterns and the cryosphere on seasonal-to-multidecadal timescales (e.g. Durack et al., 2012; Westra et al., 2014; Haumann et al., 2016; Lenderink and Van Meijgaard, 2008; Yu et al., 2021). Observational records indicate that open-ocean regions of higher salinity have become increasingly saline whilst regions of lower salinity have become fresher (i.e. an amplification of 5 %–8 % or up to ± 0.05 pss per decade since 1950; Durack et al., 2012; Zika et al., 2018), enhancing sub-surface salinity in the subtropics and freshening the tropics, sub-polar, and polar regions (Durack and Wijffels, 2010; Fox-Kemper et al., 2021). Climate models predict that this trend will be further amplified in future (Fox-Kemper et al., 2021), resulting in spatially variable changes in open-ocean sea surface salinity on the order of an increase/decrease of 0.5 to 1 pss by 2100 relative to the 1985–2014 mean (Röthig et al., 2023). Salinity in coastal areas, continental shelf seas, marginal seas, and (semi-)enclosed basins naturally fluctuates on daily-to-decadal timescales, and salinity trends are more localised and complex, related to a combination of meteorological events, climate-driven changes in the hydrological cycle, and the impact of other local anthropogenic stressors (e.g. land use, vegetation cover, and water management pressures). The complexity of factors influencing salinity in these regions contributes to uncertainty in the magnitude and direction of future salinity trends in coastal areas, marginal seas, and (semi-)enclosed basins.

Salinity is a major abiotic factor influencing marine ecosystem structure and function, but the impact of salinity on marine organisms has received relatively little attention compared to other climate-related environmental stressors (e.g. Röthig et al., 2023, and references therein). Salinity stress triggers a range of metabolic responses, including internal osmotic and ionic adjustments that can lead to morphological changes, changes to photosynthesis and respiration rates, and biochemical changes such as to osmolyte synthesis (e.g. Kirst, 1990). The response of marine phytoplankton to salinity stress has largely been investigated in coastal and euryhaline species that naturally experience variable salinity conditions, including extreme and/or short-lived salinity events. For example, at one station on the French Atlantic coast, the typical winter salinity range was 21 to 36, with on average four short-lived extreme low-salinity events lasting from a few days to a few weeks (transient salinity decreases of approx. 2 to > 8) each year (Poppeschi et al., 2021). In the open ocean, extreme salinity anomalies are more typically defined by shifts of 0.2 to 1 pss associated with freshening events; the transit of mesoscale eddies; and atmosphere–ocean dynamics related to the El Niño – Southern Oscillation, Madden–Julian Oscillation and North Atlantic Oscilla-

tion (Liu et al., 2023). Additionally, seasonal monsoons can freshen sea surface salinity by up to 8 pss for several weeks (e.g. at one Bay of Bengal mooring in 2015; Weller et al., 2019), and tropical cyclones can shift salinity by as much as 6 pss for days to weeks (H. Xu et al., 2020). Both coastal and open-ocean phytoplankton species therefore experience varying degrees of salinity change over a range of short- and long-term timescales. Laboratory experiments indicate that some marine phytoplankton species are tolerant to salinity change whilst others are not, suggesting that the impact of salinity on phytoplankton physiology may be strain specific (e.g. Brand, 1984).

One marine phytoplankton species with a demonstrated broad salinity tolerance is the coccolithophore *Emiliania huxleyi*, which can grow under salinities as low as 11–15 (Brand, 1984; Paasche et al., 1996; Lohbeck et al., 2013) and as high as 38–45 (Bukry, 1974; Brand, 1984; Winter et al., 1979; Fisher and Honjo, 1988; Gebühr et al., 2021; Linge Johnsen et al., 2019; Fielding et al., 2009). *Emiliania huxleyi* has a near-global open-ocean distribution (limited presence or absence in the very high latitudes; e.g. Winter et al., 2013) but also thrives in shelf seas and in coastal environments. For example, large blooms of *E. huxleyi* are observed in Norwegian and Patagonian coastal waters and fjords in the summer (e.g. Holligan et al., 1993; Van Der Wal et al., 1995; Winter et al., 2013; Díaz-Rosas et al., 2021), in addition to shelf sea blooms and blooms in the open ocean (e.g. Poulton et al., 2013, 2014; Tyrrell and Merico, 2004; Silkin et al., 2020).

Previous studies have demonstrated that the physiological response of coccolithophores to changing environmental conditions can influence the size of the individual plates (coccoliths) that form an inorganic calcite cell covering (coccosphere) of the coccolithophore (e.g. Faucher et al., 2020; Bollmann, 1997; Bollmann and Herrle, 2007). Laboratory experiments show that hyper- and hyposaline conditions impact the growth and morphology of many *E. huxleyi* strains (Saruwatari et al., 2016; Green et al., 1998; Paasche et al., 1996; Linge Johnsen et al., 2019; J. Xu et al., 2020; Gebühr et al., 2021). Additionally, coccolith size varies systematically along natural sea surface salinity gradients (Bollmann and Herrle, 2007; Bollmann et al., 2009). In light of this morphological sensitivity to salinity conditions, a transfer function relating *E. huxleyi* coccolith size and salinity has been developed to derive paleosalinity records independently of other geochemical proxies for use in paleoclimate and paleoceanography research (Herrle et al., 2018; Bollmann et al., 2009; Bollmann and Herrle, 2007). Cellular osmotic adjustments that change cell size have been proposed as the driver of the observed morphological response of *E. huxleyi* coccoliths to salinity stress (Bollmann et al., 2009; Gebühr et al., 2021), but the mechanistic link between salinity and cellular morphology remains unclear. Another open question is how rapidly *E. huxleyi* growth and morphology can respond to salinity changes, which could range from small salinity fluctuations (up to 1; e.g. Liu et al., 2023) over seasonal-

to-annual timescales in the open ocean to larger-magnitude salinity events (increases and decreases of 2 to > 10; e.g. Poppeschi et al., 2021; Ridgway, 2007; Weller et al., 2019; Malan et al., 2024; Grodsky et al., 2015), with a rapid onset and short duration that are more likely in coastal and shelf sea environments.

The aim of this study was to observe the short-term response of *E. huxleyi* growth, morphology, and calcification to an abrupt exposure to low- and high-salinity conditions by taking measurements of cell concentrations, coccosphere size, and coccolith size every 4 h for 6.5 d. Cultures were simultaneously transitioned from a light : dark cycle to continuous light at the onset of the salinity treatment, with the aim that continuous light conditions would desynchronise the cell division cycle by the end of the experiment. Whilst our sampling frequency aimed to identify how quickly morphology responded to salinity stress, an unintended advantage of our 4 h sampling regime was the real-time observation of the response of the cell division cycle to both salinity stress and the onset of continuous light conditions.

2 Methods

2.1 *Emiliana huxleyi* cultures

The two strains of *E. huxleyi* used in this study were selected to represent an isolate from a lower-marine-salinity regime contrasted with an isolate from a comparatively high-marine-salinity regime. Both strains are isolates from coastal regions that may realistically experience larger transient shifts in salinity compared to open-ocean regions. Strain PLYB11 (Plymouth Culture Collection, UK) is an isolate from the coastline near Bergen, Norway, with a natural seasonal salinity range of 25–33 (Paulino et al., 2018). Strain RCC1232 (Roscoff Culture Collection, France) is a coastal isolate from the bay of Villefranche-sur-Mer, France. The salinity range in this region of the northwest Mediterranean Sea is approximately 37–38 (Kapsenberg et al., 2017). Stock cultures of both strains were maintained at 15 °C under 12 h light : 12 h dark conditions at an irradiance of approx. 70 $\mu\text{mol photons m}^{-2} \text{s}^{-1}$ and a salinity of 35. Cultures were grown in sterile *f/2*-enriched artificial seawater prepared using a commercial sea salt mixture (Ulramarine; Waterlife Research Industries Ltd., UK) that was dissolved in Milli-Q water with the addition of 0.5 g L^{-1} tricaine to prevent salt precipitation during autoclaving.

2.2 Salinity experiments

Cultures of PLYB11 and RCC1232 were grown under three salinity conditions: 25 (low salinity/hyposaline treatment), 35 (control), and 45 (high salinity/hypersaline treatment). The low- and high-salinity treatments applied here are at the extremes of global open-ocean sea surface salinity, which typically ranges between 29 and 38 (Zweng et al.,

2013). However, they encompass the wider range of salinities present in some marine settings (e.g. salinities as low as 22–26 in regions of the Arctic Ocean and Baltic Sea and as high as 41 in the Red Sea; Umbert et al., 2024; Sofianos and Johns, 2017) and the magnitude of extreme salinity change that can occur in coastal regions (e.g. transient decreases in salinity of up to 10 have been reported at some stations and moorings; Poppeschi et al., 2021; Ridgway, 2007; Weller et al., 2019; Grodsky et al., 2015).

The *f/2*-enriched artificial seawater described above was diluted to the final experimental salinity by the addition of sterilised Milli-Q. To initiate each salinity experiment, triplicate 70 mL polycarbonate flasks with vented lids were prepared with 60 mL of sterile salinity 25, 35, or 45 media and directly inoculated with a small volume of stock culture (35 salinity) to achieve a start concentration of approx. 2×10^5 cells mL^{-1} . Culture flasks were sampled continuously every 4 h from the start of the experiment (0 h at 22:00 local time (LT) for PLYB11 and 00:00 LT for RCC1232) for 6.5 d (156 h) so that high-resolution temporal data could be collected to monitor how rapidly growth and morphology responded to the abrupt change in salinity conditions. At the end of the light phase preceding the start of the experiment (06:00 to 18:00 LT), light in the experimental incubator was left on continuously for the duration of the experiment, thereby switching the experimental cultures from a 12 h light : 12 h dark regime to a continuous light regime at 0 h. Continuous light conditions are often used to desynchronise the cell cycle (e.g. Müller et al., 2008, 2015), and with our 4 h measurement frequency we were able to additionally track how long the cell division cycle remained entrained to the previous 12 : 12 light : dark regime by analysing changes in cell concentrations and cell size through the experiments.

2.3 Cell concentration and growth

At each sampling time point, flasks were gently mixed to ensure the homogenous suspension of cells throughout the seawater before sampling and to re-equilibrate air in the flask headspace with sterile air. The sampling time points for cell concentration for PLYB11 were 22:00, 02:00, 06:00, 10:00, 14:00, and 18:00 LT. For strain RCC1232, the sampling time points were 00:00, 04:00, 08:00, 12:00, 16:00, and 20:00 LT. The cell concentration data at 10:00 (PLYB11) and 12:00 (RCC1232) LT (7 of a total of 40 sampling time points for each experiment, selected to represent a 24 h sampling frequency) have been previously presented in Gebühr et al. (2021).

Cell concentration was determined from a 400 μL aliquot of culture (dilution factor of 26) using an automatic cell counter (CASY Model TT; OMNI Life Science). Cell concentrations are reported as viable cells measured between 3.00 and 20.03 μm using a 60 μm capillary, the lower size threshold having been determined for these strains prior to the start of the experiment. The CASY cell counter also re-

ports particle size distribution in the sample, and here we report “cell” size from CASY derived at 4 h time intervals (coccoliths not removed from cells) from the mean particle volume measured between 3.00 and 20.03 μm on the same aliquot. Whilst we refer to CASY-derived mean particle size measurements as cell size in this study, note that the particle size reported by CASY is based on the electrical resistance of particles and therefore represents an intermediate size between cell and coccosphere size (Gerecht et al., 2015), thus overestimating true cell size but enabling accurate and rapid monitoring of the pace of fluctuation between smaller and larger mean particle sizes in the cultures for the purpose of monitoring cell division over time. For this reason, we use size measurements obtained through microscopy (Sect. 2.4) rather than CASY for accurate coccolith and coccosphere size data to investigate the short-term response of morphology and calcification to the salinity treatments applied in this study. Particle volume measurements from CASY have a maximum error of $\pm 2\%$ (OMNI Life Science).

Daily growth rates $\mu_{24\text{h}}$ (d^{-1}) were calculated from cell concentration data following Eq. (1),

$$\mu_{24\text{h}} = \frac{\ln N_t - \ln N_{t-24\text{h}}}{t - t_{-24\text{h}}}, \quad (1)$$

where $N_{t-24\text{h}}$ and N_t are the cell concentrations, N , of the culture at two sampling time points, t and $t_{-24\text{h}}$, that are consecutive but separated by 24 h, e.g. sampling at 22:00 LT or at 14:00 LT on 2 consecutive days (Guillard, 1973; Wood et al., 2005).

Instantaneous cell division rates μ_t (h^{-1}) were calculated for overlapping 8 h time intervals following Eq. (2) (Nelson and Brand, 1979).

$$\mu_t = \frac{1}{N_t} \frac{N_{t+4\text{h}} - N_{t-4\text{h}}}{8\text{h}} \quad (2)$$

2.4 Coccolith and coccosphere morphometrics

Samples for measurement of coccosphere diameter (size including the external layer(s) of coccoliths, \emptyset) and coccolith length (C_L) using scanning electron microscopy (SEM) were taken approx. every 8 h beginning at 12 h. At each sampling time point, 2 to 5 mL of culture from each flask was gently filtered on a polycarbonate filter (0.8 μm pore size, 25 mm diameter) using a borosilicate vacuum filtration flask (Millipore) and air dried for 24 h. For each salinity treatment, one of these triplicate filters was then mounted onto an aluminium stub and sputter coated with 4 nm platinum for SEM imaging using a field-emission SEM (Zeiss SIGMA VP). Measurements at 0 h were taken from a single filter from the salinity 35 stock culture used to inoculate each experimental flask (i.e. the size measurements at 0 h are the same for all salinity experiments for each strain).

Coccolith size measurements were made for a minimum of 50 flat-lying individual coccoliths at a magnification of

20 000 \times from each filter, and at least 50 intact coccospheres were imaged at a magnification of 10 000 \times for coccosphere size measurements. Measurements were made using ImageJ (v.1.51) from images with a dimension of 1024 \times 768 pixels. Coccolith and coccosphere size measurements were calibrated to 2 μm polystyrene calibration beads (certified mean diameter $1.998 \pm 0.016 \mu\text{m}$; Duke Standards Microsphere 4000 Series, certified batch number 4202–008) filtered onto a polycarbonate filter and measured once vertically and once horizontally, then averaged. Coccosphere size was similarly measured as the average of one vertical and one horizontal measurement (Lamoureux and Bollmann, 2004; Nederbragt et al., 2004).

Measurements of \emptyset , C_L , and coccolith width from a single sampling time point on experiment day 7 (14 July 2014 10:00 LT for PLYB11, all salinities) and at a single sampling time point on experiment day 5 (12 July 2014 12:00 LT salinity 25 and 35) or day 7 (13 July 2014 20:00 LT salinity 45) for RCC1232 have previously been presented in Gebühr et al. (2021). Morphometric data for the approx. 18 other sampling time points in each experiment are presented here for the first time.

2.5 Cellular particulate organic and inorganic carbon content

To further quantify the impact of morphometric responses to short-term salinity stress on calcification and cellular biogeochemical traits of *E. huxleyi*, we estimated cellular particulate inorganic carbon content (PIC), particulate organic carbon content (POC) and the inorganic to organic carbon ratio (PIC:POC) of both strains towards the end of each salinity experiment. Cellular PIC was estimated following the method of Young and Ziveri (2000), where cellular PIC is a function of C_L and the number of coccoliths per coccosphere (C_N) following Eq. (3),

$$\text{cellular PIC (pg cell}^{-1}\text{)} = \left(C_L^3 \times 2.7 \times K_s \right) \times C_N, \quad (3)$$

where K_s is a shape factor that describes species-specific coccolith morphology, and 2.7 is the density of calcite ($\text{pg } \mu\text{m}^{-3}$). Various K_s values have been published for *E. huxleyi*, but here we use K_s values published by Linge Johnsen et al. (2019; their Table 3) derived from thickness measurements under circular polarised light microscopy that are specific to strain PLYB11 (mean $K_s = 0.014$ for salinity 25 and 0.015 for salinity 35 and 45) or strain RCC1232 (mean $K_s = 0.017$ for salinity 45 and 0.019 for salinity 25 and 35) and were from experiments using the same salinity conditions as applied here. Within sample 148 h (PLYB11) and 140 h (RCC1232) from each salinity experiment, the C_N of 30 individual coccospheres (the number of coccoliths visible on each coccosphere surface multiplied by 2 to approximate the total number of coccoliths per coccosphere) and the C_L of one of the coccoliths present on the surface of the

same coccospheres (i.e. distinct from the C_L measured on exclusively loose flat-lying coccoliths) were measured from each SEM image to estimate cell-specific PIC using Eq. (3).

Cellular biomass (POC) was estimated as a function of cell volume. Cell diameter cannot be directly measured from SEM images (as the coccosphere obscures the internal organic biomass) and was therefore estimated for the same 30 coccospheres in the 148 and 140 h samples of both strains by subtracting $2 \times$ coccolith thickness (using mean coccolith thickness values for PLYB11 and RCC1232 at salinity 25, 35, and 45 conditions from Linge Johnsen et al., 2019) from our measurements of coccosphere diameter for each cell. Cell POC was then estimated following Eq. (4) (Menden-Deuer and Lessard, 2000, for prymnesiophytes),

$$\text{cellular biomass, POC (pg cell}^{-1}\text{)} = 0.23(\text{cell volume})^{0.9}. \quad (4)$$

2.6 Media chemistry

The pH and total alkalinity were measured at the beginning (0 h) and end (156 h) of each salinity experiment. A portable pH meter (WTW Multi 3400i; Xylem Analytics, Germany) was used to measure pH, and a titration method (MQuant Alkalinity Test; Merck) was used to measure alkalinity. Dissolved inorganic carbon (DIC) was approximately calculated as the difference between the two acid capacity values determined through titration ($K_{S4.3} - K_{S8.2}$). The media chemistry at the start and end of these salinity experiments was previously reported in a supplementary dataset (Table S4) accompanying Gebühr et al. (2021) and can be accessed through the online supporting information of that publication (see “Data availability” section).

2.7 Statistical analysis and data visualisation

Statistical analysis was performed using GraphPad Prism for macOS (v8.4.1, GraphPad Software, LLC). A one-way ANOVA was performed to assess statistical changes in coccolith size or coccosphere diameter through time within each salinity experiment, with Tukey’s post hoc test to identify the source of the main effect determined by ANOVA. Data were considered significant at the 95 % confidence interval ($p < 0.05$). Data figures were plotted in GraphPad Prism, and the final layout was arranged using Adobe Illustrator.

3 Results

3.1 Cell division phasing and growth under three salinity conditions

Under all salinity conditions, the mean cell size and cell concentration of *E. huxleyi* PLYB11 cultures during the experiments (note that here cell size is the particle size measured using a CASY particle counter, which is an intermediate

measurement between cell and coccosphere size and therefore an overestimate of true cell size; see Methods) show a repeating cyclicity that represents a phased cell division cycle (Fig. 1a–f); i.e. a portion of (but not all) cells in the culture are moving collectively through the cell division cycle each 24 h period (see also Chisholm, 1981, for terminology). This cell division cycle phasing persisted for the complete duration of the experiment (6.5 d) after the onset of continuous light conditions at 0 h (previously acclimated to 12 : 12 light : dark) under all salinity conditions. Phasing of the cell division cycle was identified by distinct maxima and minima in mean cell diameter that were repeated with an interval of 24 ± 4 h, as well as by regular intervals of increasing cell concentration (indicating an interval of population division) followed by a plateau in cell concentration (indicating an interval of production or no cell division). Intervals of increasing cell concentration aligned with the occurrence of cell size minima, as would be expected during an interval of phased cell division. Under all salinity conditions, consecutive cell size minima and periods of increasing cell concentration occurred during what would have been the last 4–8 h of the pre-experiment dark period (indicated by the alternating grey shading on the x axis of Fig. 1).

Cyclical fluctuations in cell concentrations and cell diameter were also apparent in strain RCC1232 under all salinities (Fig. 1g–l). However, intervals of cell concentration increase/stasis and shifts from cell size minima to cell size maxima were less uniform, and there was a smaller size change between successive cell diameter minima and maxima in this strain compared to PLYB11. Cell diameter fluctuations in RCC1232 were most pronounced under salinity 45 (Fig. 2k). Mean cell diameter minima occurred every 24 to 28 h under all salinity conditions, with longer intervals between cell diameter minima occurring after approx. 72 h of the experiment. Intervals of minimum cell diameter coincided with the end of an interval of increasing cell concentration, comparable to the observations for PLYB11.

The 156 h duration of the experiments represented just over six division–production cycles, during which approximately one-third to one-half of the population moved through cell division (growth rates, μ_{24h} , ranging from 0.26 d^{-1} in PLYB11 under salinity 45 to 0.41 d^{-1} in RCC1232 under salinity 45). The experiment duration therefore represented between two and four complete generations under the salinity conditions. Instantaneous cell division rates (μ_t , 8 h averaging) fluctuated between a minimum of -0.01 h^{-1} and 0.02 – 0.049 h^{-1} at salinity 25, 0.03 – 0.064 h^{-1} at salinity 35, and 0.015 – 0.033 h^{-1} at salinity 45 for PLYB11 (Fig. 2a–c), with an interval of 24 ± 4 h between peak division rates throughout the experiments. For RCC1232, instantaneous cell division rates (Fig. 2e–g) fluctuated between -0.01 h^{-1} (minimum for all salinity treatments) and 0.015 – 0.092 h^{-1} at salinity 25, 0.014 – 0.075 h^{-1} at salinity 35, and 0.017 – 0.044 at salinity 45, with an interval of 24 ± 4 h between peak division rates. Division rate minima occurred during what would

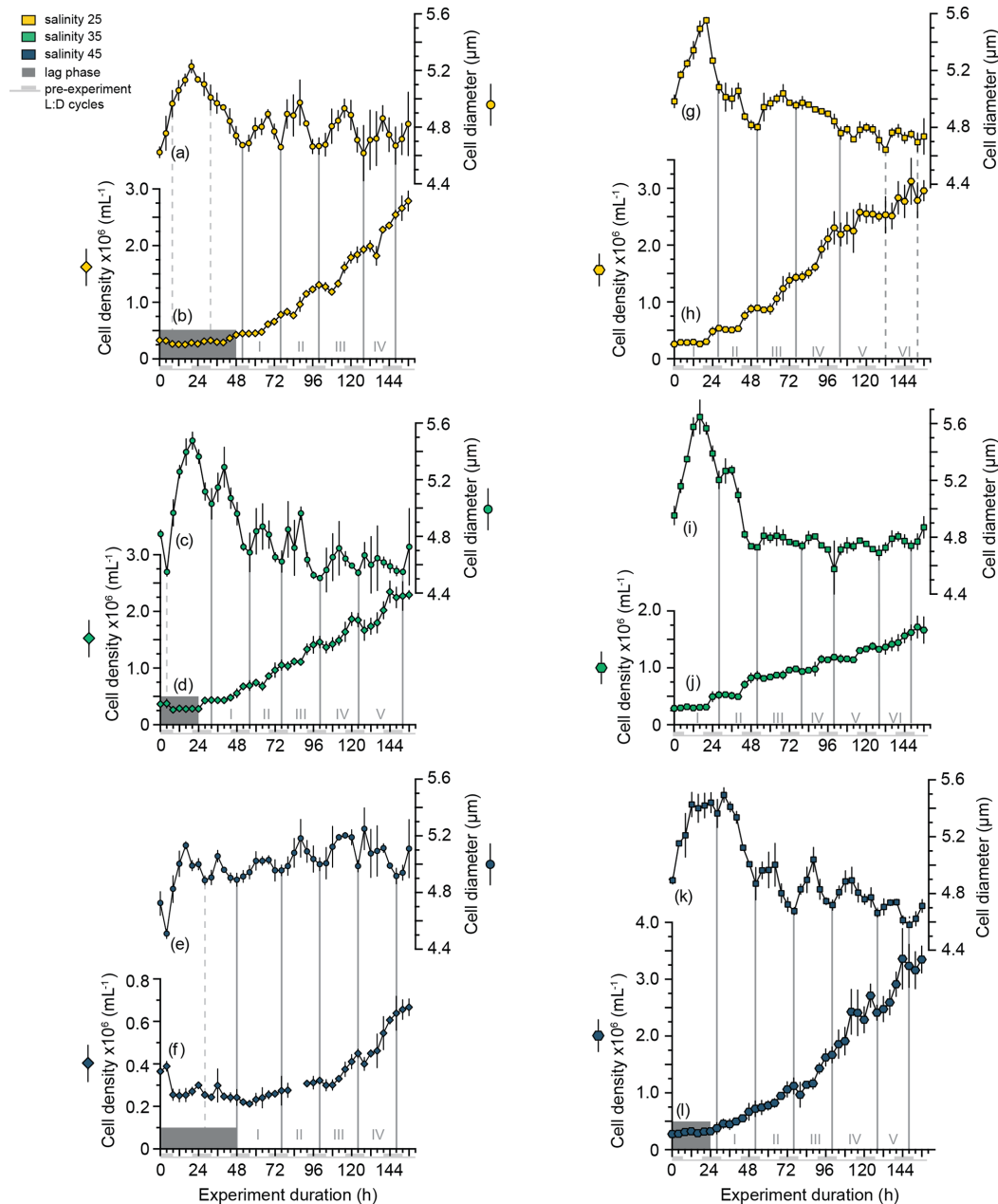


Figure 1. Cell density and cell size (derived from the CASY particle counter) of *E. huxleyi* strains PLYB11 (a–f) and RCC1232 (g–l) over the 156 h duration of experiments under salinity 25 (hyposaline), salinity 35 (control), and salinity 45 (hypersaline) conditions. Each data point represents the mean and standard deviation of triplicate measurements. Note the different y-axis scale in (f). Sequential population division cycles are indicated by the vertical grey lines and numbered with Roman numerals, determined by mean cell size minima. Some experiments exhibit a period of minimal growth or an initial lag phase in the first 24–48 h, indicated by the dark-grey shaded area. Inferred division cycles that occur during these initial lag phases are denoted by dashed grey lines. For context, the timing of pre-experiment alternations between light and dark (L : D) conditions are shown as light grey shading on the *x*-axis labels (note that continuous light conditions were applied for the duration of all experiments; see Methods).

have been the last 4–8 h of the pre-experiment dark period or the first 4–8 h of the pre-experiment light period in both strains. In the first 48 h, RCC1232 under salinity 25 and salinity 35 conditions shows large peak division rates that decrease by more than half by 72 h. Such a large change in

peak division rates between the first 2 d and the remainder of the experiment duration is not observed in PLYB11. After 3–4 d of growth under the experimental treatments (between 80 and 152 h), periodicity in cell division rates clearly persists (Fig. 2d and h), but peak division rates start to de-

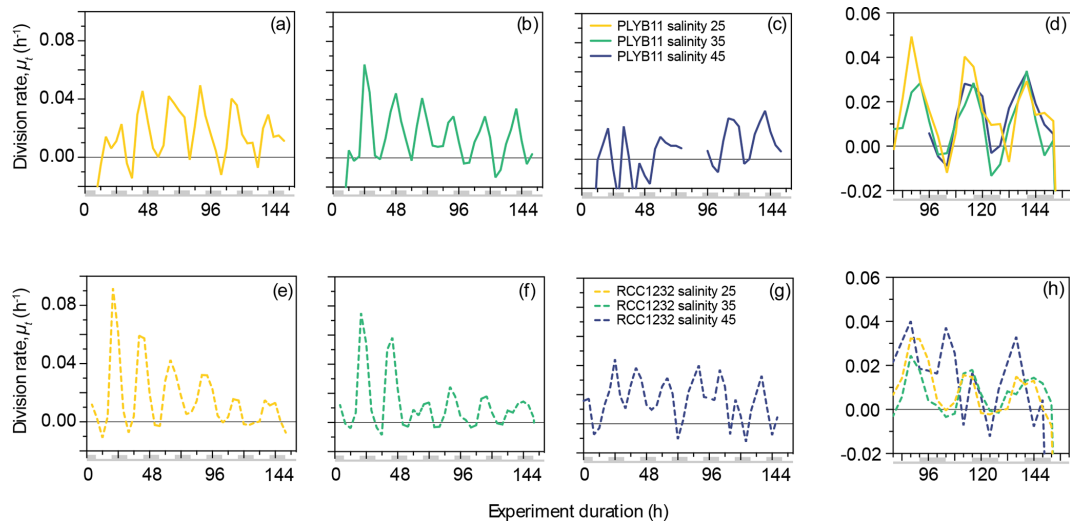


Figure 2. Division rates, μ_t , of *E. huxleyi* strains PLYB11 (a–d) and RCC1232 (e–h) over the duration of experiments under salinity 25 (hyposaline), salinity 35 (control), and salinity 45 (hypersaline) conditions. For comparison across the three salinity treatments, division rates over days 4–6.5 of the experiments are shown in (d) for PLYB11 and in (h) for RCC1232 (note the different y-axis scale). Similarly to Fig. 1, the timing of pre-experiment alternations between light and dark conditions are shown as light-grey shading on the x-axis labels for context.

cline in both strains under salinity 25 and 35 by approx. 144 h (peak μ_t decreasing from 0.04–0.06 h^{-1} to 0.02–0.05 h^{-1} in PLYB11 and peak μ_t decreasing from 0.025–0.035 h^{-1} to 0.01–0.015 h^{-1} in RCC1232). In days 4–6 of the salinity 45 experiment with RCC1232, a 4–8 h offset between the timing of peak division and the timing of peak division under salinity 25 and 35 also emerges (Fig. 2h). For PLYB11, peak division rates in the final 3 d of each salinity experiment all occur at the same time point (Fig. 2d).

Based on the logarithmic transformation of cell concentration data, all PLYB11 experiments (including control salinity 35) and the RCC1232 salinity 45 experiment showed an initial lag phase of 24 to 48 h. Mean cell size increased by 0.5–0.9 μm over the first approx. 12 h of each experiment before decreasing again over the following 12–24 h. This was equivalent to a mean cell size increase of 11 % (RCC1232) or 13 % (PLYB11) at salinity 25 and 45 and an increase of 14 % (RCC1232) or 20 % (PLYB11) at salinity 35 (Figs. 1 and 2). In the RCC1232 and PLYB11 salinity 35 and 45 experiments, the subsequent mean cell size decrease was not as great as the initial size increase in the first 12–24 h of the experiment and was followed by a smaller size increase at the beginning of division cycle II (similar in magnitude to subsequent mean size minima–maxima fluctuations for the remainder of the experiment). In both strains under salinity 35, division cycle II (between 32 and 56 h in PLYB11 and 28 and 52 h in RCC1232) saw a large size decrease (approx. –10 %) during the interval of increasing cell concentrations.

3.2 Effect of salinity on coccolith and cell size in PLYB11

Under control salinity 35 conditions, mean coccolith size (C_L) and coccosphere size (\varnothing) in PLYB11 remained relatively unchanged for the duration of the experiment, fluctuating between 2.6–2.9 μm and 4.6–5.2 μm , respectively, but with no consistent temporal trend (Fig. 3b and e). Following the abrupt transition to salinity 25, mean C_L responded with a significant decrease from 2.9 to 2.6 μm (–10 %) within the first 12 h of exposure (one-way ANOVA, $F(7,410) = 6.493$, $p < 0.0001$; the Tukey post hoc test showed that C_L measured at 0 h is statistically larger than all other measured time points and no other multiple comparisons were statistically significant, Fig. 3a) and mean C_L remained at 2.5–2.7 μm for the remainder of the salinity 25 experiment. Under salinity 45, mean C_L increased in two stages (Fig. 3c), with a small but significant step increase of 7 % at 36 h and a further 9 % increase at 100 h. Mean \varnothing also changed at a similar time point to C_L change under salinity 45 (Fig. 3d–f), gradually increasing from 4.9 μm at 76 h to 5.3 μm at 100 h (an equivalent mean cell volume increase of 27 %), after which mean \varnothing remained relatively constant at 5.1–5.3 μm (Fig. 3f). In contrast, the response of mean \varnothing to salinity 25 occurred later in the experiment than the response of mean C_L , decreasing from 5.0 to 4.5 μm between 100 and 132 h. The mean \varnothing at the end of the salinity 25 experiment (156 h) was 4.2 μm (an equivalent cell volume decrease of 41 %).

As phasing of the cell division cycle persisted under continuous light for the entire duration of the experiment, the analysis of the short-term effect of salinity on coccolith size

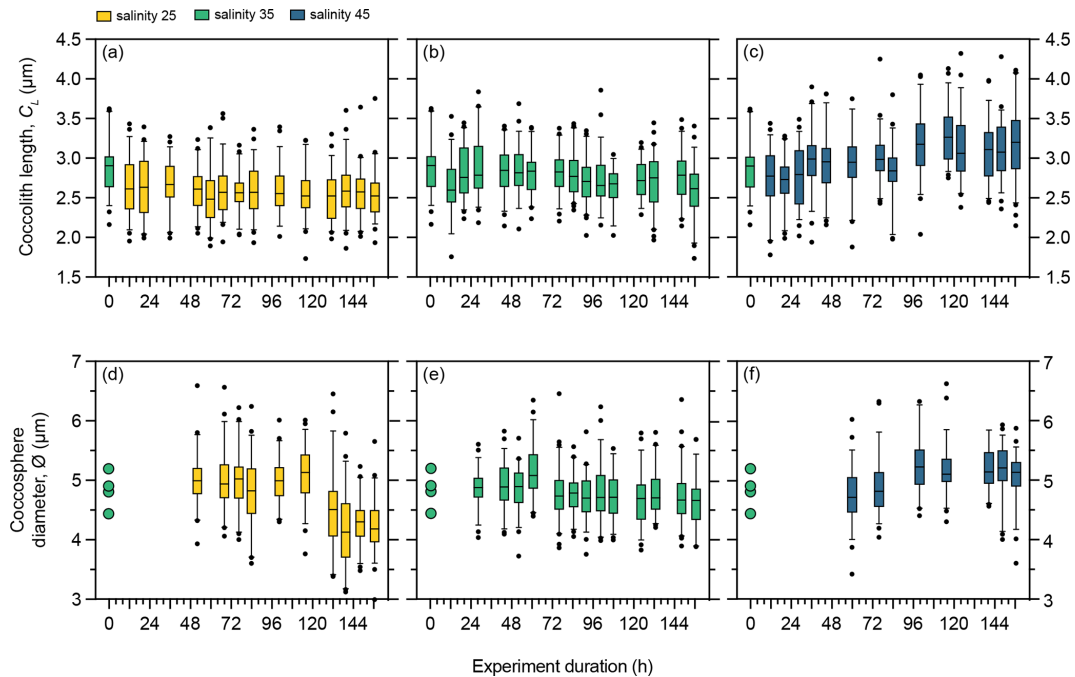


Figure 3. Morphology of *E. huxleyi* strain PLYB11 under three salinity conditions over the 156 h exposure duration. (a–c) Coccolith length under salinity 25, salinity 35, and salinity 45. (d–f) Coccosphere diameter under salinity 25, salinity 35, and salinity 45. Box and whisker plots show the 25th–75th quartiles (box with the median shown by the central line) and 5th–95th quartiles (whiskers) of a minimum of 50 measurements at each time point. Measurements outside the 5th–95th quartiles of the data are shown as circles. Measurements at time 0 h are taken from the stock control culture (salinity 35) used to inoculate all flasks and are therefore also shown as the start coccolith length or coccosphere diameter measurements for the salinity 25 and 45 experiments at 0 h. The 0 h sample for PLYB11 coccosphere diameter measurements was lost, but a small number of coccosphere measurements ($n = 4$) could be made from coccospheres imaged within the coccolith length sample and are provided (circles, **d** and **e**) to indicate coccosphere diameter at 0 h in this experiment.

and cell size must be based on samples taken at the same time point within the cell cycle to avoid comparing data from later in the production phase (when cells are larger) with data from the division phase or early production phase (when cells are smaller as they have recently divided). We therefore compared temporal changes in coccolith and coccosphere size of PLYB11 under exposure to salinity 25 and 45 based on measurements taken from the same cell cycle point (cell size minima) at 76, 100, 124, and 148 h to see if the magnitude and/or direction of salinity effects on morphology remained constant with time or changed through time as the population was exposed to the new salinity condition for longer (Fig. 4). After just 3 d of growth (76 h), there was a clear difference in PLYB11 C_L between low-salinity, control, and high-salinity conditions, with larger C_L under higher-salinity conditions (Fig. 4a). The size difference between salinity 35 and salinity 45 coccoliths becomes even more pronounced by days 4–6, i.e. with longer exposure to hypersaline conditions. By contrast, the effect of salinity on \varnothing develops more steadily as the experiment progresses (Fig. 4b): after 3–4 d growth (76 to 100 h), coccospheres tend to be larger at salinity 25 and 45 than at salinity 35; however, after 6 d of growth (148 h), mean \varnothing is smallest at salinity 25 and largest at salinity 45.

3.3 Effect of salinity on coccolith and cell size in RCC1232

In contrast to PLYB11 under control conditions, RCC1232 grown under salinity 35 showed an increase in C_L within the first 48 h (Fig. 5b; one-way ANOVA, $F(13\ 744) = 4.316$, $p < 0.0001$; the Tukey post hoc test showed that C_L measured at 12 h was significantly smaller than all other time points except 84, 108, and 116 h, and no other multiple comparisons were statistically significant). A significant increase in \varnothing within the first 24 h in the salinity 35 experiment is also clear (Fig. 5e, one-way ANOVA, $F(11\ 647) = 14.59$, $p < 0.0001$; the Tukey post hoc test showed that C_L measured at 0 h is statistically smaller than all other measured time points) and persisted for the remainder of the experiment (Fig. 5b and e).

The Mediterranean strain RCC1232 showed a negligible C_L response to salinity 25 relative to C_L at 0 h, but C_L increased by approx. 18 % under salinity 45 conditions within the first 36 h of the experiment (Fig. 5a and c). A significant 28 % increase in \varnothing is recorded between 0 and 36 h under salinity 25 (Fig. 5d; Tukey multiple comparison test, $p < 0.0001$, 95 % CI = -1.550 to -0.9713). Intact coccospheres were rarely observed on filters after 72 h growth un-

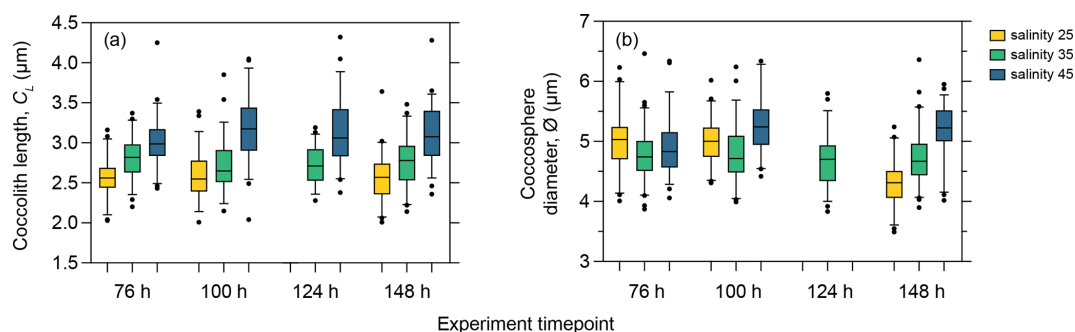


Figure 4. Coccolith length (a) and coccosphere diameter (b) response of *E. huxleyi* strain PLYB11 after 76 h (approx. 3 d), 100 h (approx. 4 d), 124 h (approx. 5 d), and 148 h (approx. 6 d) exposure to salinity 25, salinity 35, and salinity 45 conditions. The measurements from these selected time points are sampled from the same cell cycle time point (cell size minima). Samples for \varnothing were not taken at 124 h in the salinity 25 and salinity 45 experiments.

der low salinity (Fig. S1 of Gebühr et al., 2024; see “Data availability” section). RCC1232 mean \varnothing increased by 27 % over the course of the salinity 45 experiment, but the \varnothing increase primarily occurred 2 d earlier, within the first 48 h (Fig. 5f).

To assess the overall impact of low- and high-salinity treatments on C_L and \varnothing , measurements from the same points of the cell division cycle were shown for PLYB11 in Fig. 4. A comparable analysis was not possible for RCC1232 because fewer overall SEM measurements of \varnothing were taken (and were not possible under salinity 25, as explained above) and because the time points of C_L SEM measurements unfortunately did not align well with the time points of cell size minima/maxima as determined from CASY data (Fig. 1g–l). However, after 3–4 d of growth under each salinity condition, RCC1232 C_L was consistently smallest under salinity 25 and largest under salinity 45 (Fig. 5). RCC1232 \varnothing was less variable across the different salinity conditions than was observed in PLYB11 (based on SEM measurements; Figs. 3–5) but also appears to have not completely stabilised after > 6 d under the new salinity conditions (based on CASY measurements; Fig. 1). Under all salinity conditions, size measured at 0 h from the control 35 salinity inoculum was also substantially smaller than \varnothing measured at all other time points (Fig. 5).

3.4 Calcification response to salinity

Our short-term experiments show that the rapid effect of salinity on *E. huxleyi* morphology leads to changes in the inorganic carbon per coccolith and per cell of both strains between each salinity condition (Table 1). After approx. 6 d growth, the difference in the size of PLYB11 coccoliths between salinity 25 and salinity 45 conditions (+22 %) in PLYB11 is equivalent to a 95 % increase in coccolith particulate inorganic carbon (PIC) compared to PIC under salinity 25 (Table 1). In RCC1232, mean coccolith PIC was 26 % larger under salinity 45 conditions relative to salinity 25 con-

ditions after approx. 6 d growth, but coccolith PIC was conversely largest at salinity 35 as mean C_L was only 0.03 μm smaller at salinity 35 than salinity 45, but the mean K_s value of RCC1232 at salinity 35 compared to salinity 45 is 12 % larger (Linge Johnsen et al., 2019). At the cellular level, PIC per coccosphere after approx. 6 d growth under salinity 45 is almost double that at salinity 25 in PLYB11 (an increase of 1.38 pgC cell^{-1} or 99 %), principally driven by the response of C_L to salinity as the number of coccoliths per cell (C_N) remains relatively unchanged between salinity conditions. Cellular PIC could not be estimated for RCC1232 under salinity 25 conditions as coccospheres were too poorly preserved (see Fig. S1 of Gebühr et al., 2024, “Data Availability” section). Cellular PIC was 4 % larger under salinity 45 compared to salinity 35 in RCC1232 after approx. 6 d growth. Larger coccosphere (cell) size with increased salinity additionally translates to a 55 % increase in cell biomass (particulate organic carbon, POC) in PLYB11 between salinity 25 and 45 conditions. The interaction between increased C_L and \varnothing with salinity and variability in C_N between individual coccospheres results in smaller differences in cellular PIC : POC between salinity treatments, although PIC : POC is slightly higher (approx. 8 %) under salinity 45 conditions.

4 Discussion

4.1 Persistent phased cell division under continuous light

Many phytoplankton exhibit cell division cycles that are synchronised (with the population doubling each day) or phased (where population doubling time exceeds 1 d) to external light : dark fluctuations (Chisholm and Brand, 1981, and references therein). Populations entrained to the light : dark cycle typically restrict cell division to a portion of the light : dark cycle (e.g. Nelson and Brand, 1979), usually the dark phase, and production occurs during the light phase (Harding et al., 1981). Cell division in *E. huxleyi* occurs pri-

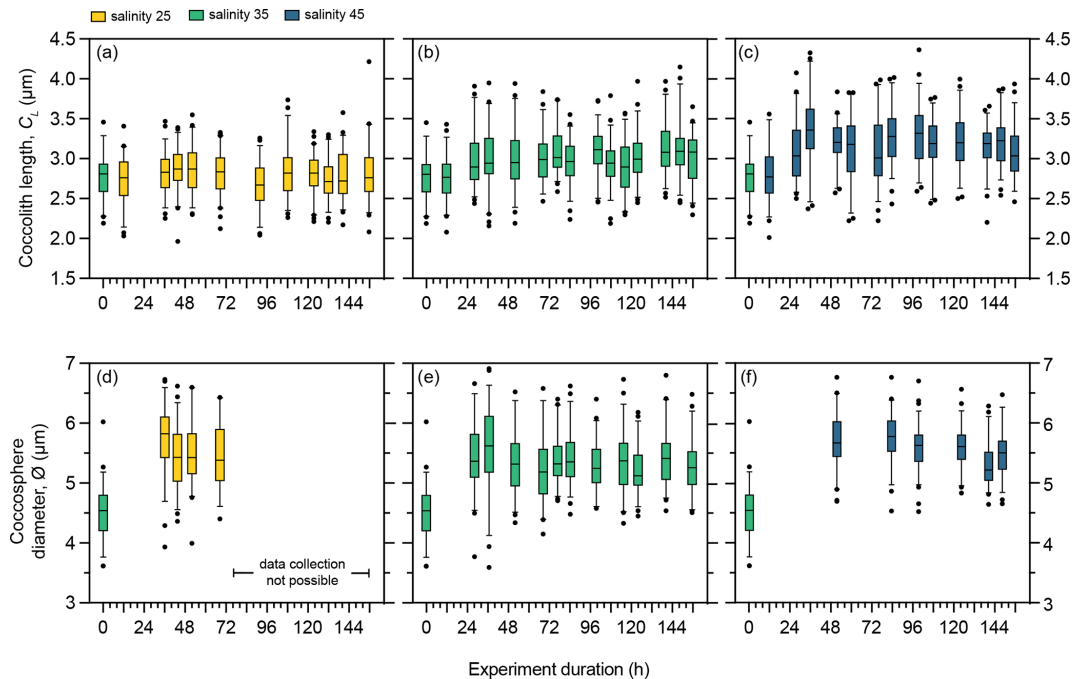


Figure 5. Morphology of *E. huxleyi* strain RCC1232 under three salinity conditions over a 156 h exposure duration. (a–c) Coccolith length (C_L) under salinity 25, salinity 35, and salinity 45. (d–f) Coccospere diameter (\emptyset) under salinity 25, salinity 35, and salinity 45. Box and whisker plots show the 25th–75th quartiles (box with the median shown by the central line) and 5th–95th quartiles (whiskers) of a minimum of 50 measurements at each time point. Measurements outside the 5th–95th quartiles of the data are shown as circles. Measurements at 0 h are taken from the stock control culture (salinity 35) used to inoculate all flasks and are therefore also shown as the start C_L or \emptyset measurements for the salinity 25 and 45 experiments at 0 h. Coccospere measurements after 96 h were not possible for the salinity 25 experiment as severe coccolith malformation led to all coccospere collapsing when filtered.

marily during the dark phase (Bucciarelli et al., 2007; Müller et al., 2008) under a range of light : dark conditions (Paasche, 1967). This phased cell division leads to diel variability in parameters that are commonly measured to track physiological responses to changing environmental conditions (Kottmeier et al., 2020), such as cellular biomass (POC). Cell concentrations through time (< 24 h measurement frequency) will similarly show step-like increases (Kottmeier et al., 2020) when population division is phased, rather than the constant exponential increase in cell concentrations expected for fully desynchronised populations (Müller et al., 2015). When sampling a phased population over a period of time, measurement time points therefore need to be carefully selected so that they target the same point of the cell division cycle each 24 h period (Barcelos e Ramos et al., 2010; Kottmeier et al., 2020). This is particularly important when aiming to quantify the impact of environmental perturbations on cell physiology, as absolute increases or decreases in cellular elemental content will vary throughout the day, and comparing data collected from different time points in the cell division cycle across experiments could change the conclusions of the experiment (Kottmeier et al., 2020).

To circumvent this complexity, some experimental designs apply continuous 24 h light to cultures with the aim of fully

desynchronising the cell division cycle. For a fully desynchronised population growing under continuous light, the sampling time has no influence on measurement values as all measurement time points are representative of the daily mean production rate, cellular elemental content, or other physiological measures of interest (Jochem and Meyerdierks, 1999; Shi et al., 2009; Müller et al., 2008, 2015; Kottmeier et al., 2020). Some phytoplankton species grow poorly or not at all under continuous light (Paasche, 1967; Brand and Guillard, 1981), reportedly including the coccolithophore species *Calcidiscus leptoporus* (Brand and Guillard, 1981), the prymnesiophytes *Isochrysis galbana* and *Chrysochromulina* sp., and the holococcolith form of *Coccolithus pelagicus* (Price et al., 1998). However, several coccolithophore species, including *E. huxleyi*, have been shown to grow well under continuous light (Brand and Guillard, 1981; Price et al., 1998). Additionally, *E. huxleyi* has reportedly been maintained in culture for prolonged periods of time (i.e. several months) under continuous light (Shi et al., 2009; Müller et al., 2017), and several publications explicitly report that the cell cycle of *E. huxleyi* became desynchronised when grown under continuous light (Müller et al., 2008, 2017, 2015, 2012).

We applied continuous light conditions from the onset of our experiments (an abrupt shift from a 12 : 12 light : dark cy-

cle to continuous light at 0 h), and our 4 h sampling regime enabled the pattern of cell division to be monitored under control, low-, and high-salinity conditions for the duration of each experiment. Unexpectedly, clear cell division phasing persisted under all salinity conditions in both *E. huxleyi* strains for the entire duration of the experiments (> 6 d representing 2–4 generations; Figs. 1 and 2). If the onset of continuous light had desynchronised cell division, we would expect to see \emptyset stabilise around a constant value and continuously increasing cell concentrations through the course of the experiment (Müller et al., 2015). Instead, we see clear mean size minima and maxima in all experiments for both strains, corresponding to step-like increases and plateaus in cell concentrations that correspond closely to the timings of the light : dark alternations that preceded the onset of the experiments (Fig. 1). Towards the end of the experiments, declining maximum division rates under salinity 25 and 45 in PLYB11 (Fig. 2) may indicate that phased division of the population was starting to weaken in this strain under low- and high-salinity stress. However, it is not clear for how many more days phased division might have persisted beyond the end of the experiment. Declining division rates at the end of an experiment may also be an indicator that phased cell division continued but population growth was entering the early stationary phase; i.e. growth could no longer proceed exponentially due to one or more changes in the physiochemical conditions of the experiment as cell density increased through the duration of the experiment. Cell concentrations were dense towards the end of all experiments ($0.7\text{--}2.8 \times 10^6$ cells mL $^{-1}$ in PLYB11, $1.7\text{--}3.3 \times 10^6$ cells mL $^{-1}$ in RCC1232; Fig. 1), which resulted in a DIC increase of up to 18 % in PLYB11 experiments and a DIC decrease of up to 18 % in RCC1232 experiments by the end of the experiments relative to starting media chemistry (Gebühr et al., 2021). The effect of cell density on media chemistry, nutrient availability, and/or light environment may have been sufficient to initiate a departure from exponential growth by the end of the 6.5 d experiment duration. This is indicated under all experimental conditions for RCC1232, where growth rates calculated from cell concentration data at 24 h intervals suggest non-exponential growth after approximately 5 d (120 h in salinity 25 and 35 experiments) or 6 d (144 h under salinity 45) of experiment duration (Gebühr et al., 2021). Mean daily growth rates indicate that PLYB11 maintained exponential growth throughout the duration of the experiments (Gebühr et al., 2021).

Persistent cell cycle phasing for 3 d after transitioning from a light : dark cycle regime (14 : 10 L : D) to continuous light has previously been shown for two species of *E. huxleyi* (with a division rate of approx. 1 d $^{-1}$), the coccolithophore *Chrysothilia carterae* (previously *Hymenomonas carterae*), and three other marine phytoplankton species (Chisholm and Brand, 1981). Cell division phasing of *E. huxleyi* strain CCMP 371 was reportedly desynchronised by “...illumination over several generations with continuous light...”

(Müller et al., 2008), but the authors did not further clarify how long this took to achieve. Where publications report the use of continuous light, the pre-experiment acclimation period is stated to be between approx. 7 and 20 generations of growth (Zondervan et al., 2001, 2002; Müller et al., 2008; Bretherton et al., 2019). Assuming that these acclimation periods were sufficient to desynchronise the cell division cycle, we conclude that phasing of cell division to the pre-experiment light : dark conditions in *E. huxleyi* must persist for a minimum of 3–4 generations (as shown by Chisholm and Brand, 1981, for populations dividing approximately daily and our experiments with lower growth rates) up to approx. 15 generations or longer. This would equate to 2–3 weeks of growth under experimental conditions for populations dividing with a growth rate of 0.7 d $^{-1}$ and approx. 4–5 weeks for populations dividing with a growth rate of 0.35 d $^{-1}$. Interestingly, cell cycle entrainment to light : dark cycles may be highly persistent in some strains or in combination with certain environmental stressors, as a previous publication reports culturing *E. huxleyi* for 6 months under continuous light to ensure complete desynchronisation of the division cycle (Müller et al., 2017).

4.2 Growth and morphological responses to continuous light

Many phytoplankton, including coccolithophores, have higher growth rates under higher irradiance levels, under longer day lengths, or when continuous light is used (e.g. Chisholm and Brand, 1981; Harris et al., 2009; Sheward et al., 2023). We did not measure growth rate before the start of the experiment so are unable to quantify the impact of the abrupt transition from a 12 : 12 L : D cycle to continuous light on growth rate under control conditions. However, both strains showed an initial phase of low or negligible increase in cell concentrations over the first approx. 24 h in the control experiment, suggesting that an immediate physiological response to the onset of continuous light probably occurred in all experiments. There are conflicting reports as to whether *E. huxleyi* has higher (Chisholm and Brand, 1981; Price et al., 1998; Bretherton et al., 2019), lower (Van Rijssel and Gieskes, 2002), or comparable (Zondervan et al., 2001, 2002; Rost et al., 2002; Nielsen, 1997) growth rates under continuous light compared to a light : dark regime. The effect of continuous or discontinuous irradiance on *E. huxleyi* growth rate is likely to be strain specific (Price et al., 1998; Bretherton et al., 2019) and/or vary depending on the combination of irradiance level and day length used (Paasche, 1967; Rost et al., 2002). Subsequently, we cannot rule out that a growth rate response to the abrupt onset of continuous light contributed to growth rate differences between low- and high-salinity conditions over the course of our experiments. However, we note that Rost et al. (2002) report that growth rates of the same strain of *E. huxleyi* (PLYB91/11) are comparable under both

a 16 : 10 L : D cycle and continuous light when grown under similar irradiance levels (70–100 $\mu\text{mol photons m}^{-2} \text{s}^{-1}$).

In the RCC1232 control experiment, both C_L and \varnothing increase within approx. 28 h of inoculation and then remain at these larger mean values (with some fluctuations) for the remainder of the experiment (Fig. 5b and e). When combined with the overall C_L and \varnothing response to low- and high-salinity conditions over the course of the experiment, this unexpected increase in C_L and \varnothing under control conditions strongly influences the overall response of C_L and \varnothing across salinities 25, 35, and 45 by 156 h in this strain: C_L increases from salinity 25 to salinity 45, whereas \varnothing is broadly comparable across all salinity treatments as \varnothing increased similarly (Fig. 5). The initial shift in C_L and \varnothing is unlikely to be caused by measurements taken at different points in the cell division cycle (Fig. 1). Instead, the onset of continuous light may have driven a rapid morphological response in RCC1232 that is not observed in PLYB11. Larger cell sizes have previously been reported within 5 h of exposure to higher light conditions for a non-calcifying strain of *E. huxleyi* (Darroch et al., 2015), but in contrast, no significant cell size difference was reported between continuous light and 14 : 10 L : D experiments for at least one *E. huxleyi* strain (Price et al., 1998).

4.3 Rapid morphological responses to abrupt salinity stress

The physiology, cellular composition, and gene expression of *E. huxleyi* can respond rapidly to abrupt and short-term (hours to days) changes in carbonate chemistry, light environment, and nutrient levels (e.g. Barcelos e Ramos et al., 2010, 2012; Iglesias-Rodriguez et al., 2017; Darroch et al., 2015). Experiments with other phytoplankton groups have demonstrated that sudden salinity perturbations induce a cascade of rapid metabolic responses within cells, some of which may be coupled to changes in cell size. For example, it only takes seconds to minutes for the green halophilic algae *Dunaliella* to adjust to osmotic differences between the external medium and the cell cytoplasm through rapid, passive water efflux or influx (Weiss and Pick, 1990), which simultaneously drives changes in cell size and volume (Maeda and Thompson, 1986). The impacts of this passive osmotic adjustment on cell size persist for minutes to hours (Weiss and Pick, 1990; Maeda and Thompson, 1986) depending on how rapidly *Dunaliella* restores its ionic equilibrium, e.g. through regulating glycol metabolism (Borowitzka, 2018). To date, the short-term (hours to days) sequence of metabolic responses of *E. huxleyi* under salinity stress remains unknown. Whilst our 4 h sampling frequency is insufficient to capture cell size changes occurring due to turgor pressure adjustment within the first seconds to minutes of exposure to hyposaline (salinity 25) and hypersaline (salinity 45) conditions, as were observed in *Dunaliella*, our experiments do capture rapid changes in *E. huxleyi* growth and morphology over the hours to days following the onset of salinity stress.

Both strain PLYB11 and strain RCC1232 exhibit type A coccolith morphologies (Fig. S1 of Gebühr et al., 2024, see “Data Availability” section; Young et al., 2003) but showed strain-specific differences in morphological characteristics under control conditions (see Gebühr et al., 2021, for a detailed morphological description of both strains), notably that RCC1232 coccolith and coccosphere sizes were on average 15 % larger than PLYB11 coccolith and coccosphere sizes. The timescale of \varnothing and C_L response to abrupt salinity stress (relative to initial strain-specific \varnothing and C_L , Figs. 3 and 5; see also Gebühr et al., 2021) was varied and did not show a consistent relationship to strain, salinity treatment, or morphological parameter. This is perhaps not surprising, as the response of cell/coccosphere size to salinity in *E. huxleyi* (Saruwatari et al., 2016; J. Xu et al., 2020; Gebühr et al., 2021; Hermoso and Lecasble, 2018) and in closely related *Gephyrocapsa* species (Hermoso and Lecasble, 2018) seems to be strain-specific (Hermoso and Lecasble, 2018; Gebühr et al., 2021). In addition to differences in the magnitude of cell/coccosphere size response to salinity 25 and 45 conditions between the two strains (Figs. 3–5; Table 1), Norwegian strain PLYB11 generally showed morphological changes after a longer period of exposure (96–120 h) to salinity 25 and 45 conditions compared to Mediterranean strain RCC1232 (which showed \varnothing changes within 28–36 h). As the concentration of cells in each experiment was high (10^5 to 10^6 cells mL^{-1}), by the end of each experiment (Fig. 1), we cannot rule out that changes in media chemistry, nutrient availability, or light environment may have additionally influenced coccolith length and/or coccosphere diameter. However, there is no systematic relationship between coccolith length and cell density; changes in coccolith and coccosphere size were already detectable within 2 d of abrupt salinity stress (when cell density was lower), and no statistically significant change in C_L was observed in PLYB11 under control (salinity 35) conditions. This strongly supports an interpretation that abrupt salinity stress rather than the potential impact of cell density on experimental condition was the primary driver of morphological changes in the experiments.

As cell size responds to the regulation of cell turgor pressure in phytoplankton and other plants (e.g. Kirst, 1990, and references therein), it is plausible that different timescales of morphological response are related to different rates of osmotic adjustment between strains or through regulation of osmolyte synthesis, active transport mechanisms, and/or membrane pumps under different salinity conditions. Species-specific synthesis of osmolytes and morphological responses to salinity stress have, for example, been reported for marine diatoms (Helliwell et al., 2021, and references therein) but are currently largely unknown for coccolithophores. *E. huxleyi* is a recognised producer of dimethylsulfoniopropionate (DMSP), a compatible solute that contributes to cellular osmotic balance (Kirst, 1996). Cellular concentrations of DMSP are coupled to salinity in many phytoplankton (e.g. Keller and Korjeff-Bellows, 1996; Kirst, 1996; Stefels, 2000;

Table 1. Mean (\pm SD) values for morphology and cellular biogeochemical parameters of *E. huxleyi* after approx. 6 d growth under three salinity conditions. Growth rate (μ_{24h}) is calculated between experiment day 2 and day 5 (see Methods). Morphometric data are from 148 h for PLYB11 and 140 h for RCC1232 (i.e. from comparable points of the cell division cycle, approximately a minimum in mean cell size). Coccolith length (C_L) and coccosphere diameter (\varnothing) are directly measured from SEM images for a minimum of 50 individuals. See Methods for the calculation of PIC (particulate inorganic carbon) following Young and Ziveri (2000) and POC (particulate organic carbon) following Menden-Deuer and Lessard (2000), based on morphometric measurements from 30 individual coccospheres per sample.

	C_L (μm)	\varnothing (μm)	PIC (pg C)		POC (pg C per cell)	PIC : POC (mol : mol)
			Per coccolith	Per coccosphere		
PLYB11						
Salinity 25	2.54 (0.30)	4.30 (0.39)	0.077 (0.03)	1.40 (0.43)	6.14 (1.79)	0.25 (0.10)
Salinity 35	2.76 (0.31)	4.71 (0.45)	0.110 (0.035)	1.95 (0.72)	8.00 (2.64)	0.25 (0.06)
Salinity 45	3.11 (0.36)	5.20 (0.43)	0.150 (0.054)	2.78 (0.74)	9.54 (1.93)	0.27 (0.05)
RCC1232						
Salinity 25	2.79 (0.31)	–	0.138 (0.22)	–	–	–
Salinity 35	3.11 (0.32)	5.41 (0.45)	0.191 (0.06)	3.28 (1.06)	11.84 (3.18)	0.30 (0.09)
Salinity 45	3.14 (0.29)	5.30 (0.37)	0.174 (0.04)	3.40 (1.07)	11.04 (2.28)	0.32 (0.10)

Dickson and Kirst, 1987) and in the macroalgae *Ulva* (Van Alstyne et al., 2003). Intracellular concentrations of DMSP have been shown to correlate with salinity in one open-ocean strain of *E. huxleyi* (McParland et al., 2020), in *E. huxleyi* strains from a range of environments (Fielding, 2010), and in the coccolithophore species *Gephyrocapsa oceanica* (Larsen and Beardall, 2023) and *Chrysotila carterae* (Vairavamurthy et al., 1985). Cellular DMSP also responds rapidly to some environmental stressors, within as little as 4 h under elevated light (Darroch et al., 2015), and the elevated DMSP content of Prymnesiophyceae relative to other phytoplankton groups (Keller et al., 1989; McParland and Levine, 2019) may provide an initial reserve to better tolerate rapid-onset salinity stress (Kirst, 1996). However, further investigation is needed to identify the rate of change in cellular ion and osmolyte concentrations for *E. huxleyi* under a range of salinity conditions and strain-specific strategies for osmotic adjustment, as both may impact the capacity of *E. huxleyi* to respond rapidly to the onset of salinity stress.

Changes in C_L with salinity have previously been attributed to changes in the size of the coccolith vesicle proportional to cell volume increases/decreases that occur when water influx/efflux is used to maintain cell turgor pressure (Bollmann et al., 2009; Gebühr et al., 2021). Particularly within short-term experiments as applied here, this hypothesis implies four things: (1) that \varnothing response must precede a C_L response; (2) that the size change in C_L and \varnothing must be positively correlated over some reasonable time frame (Suchéras-Marx et al., 2022) – i.e. cell size does not increase whilst coccolith size decreases or vice versa; (3) that the relative magnitude of the cell size change and the corresponding coccolith size change are reasonably proportional (i.e. a small cell size change does not drive a disproportionately large change in coccolith size); and (4) that a change in \varnothing due to

one or more mechanisms to maintain cellular homeostasis must persist for a sufficient amount of time for a corresponding increase in C_L to be quantifiable (relative to coccolith production rates).

Overall, our data are consistent with these criteria (Figs. 3 and 5). However, we do see intervals within each experiment where large step-like changes in \varnothing do not correspond proportionally to changes in C_L over the following hours to days (e.g. both strains under salinity 25; Figs. 3 and 5). In some cases (e.g. both strains under salinity 45) we also see \varnothing and C_L changes that occur relatively synchronously (although there is some discrepancy in the C_L and \varnothing sampling time points). Generally, changes in C_L tend to emerge more gradually and after a longer period of exposure than changes in \varnothing (Figs. 3 and 5), apart from RCC1232 under salinity 45 where the C_L increases between 24 and 36 h occur within the same time frame as the \varnothing increase. The measured loose coccoliths initially represent a mix of coccoliths produced during pre-experiment conditions and coccoliths produced under salinity stress, with a diminishing contribution from pre-experiment coccoliths to C_L measurements as the experiment progresses. New coccoliths are produced at least every approx. 60 min (Paasche, 2002; Suchéras-Marx et al., 2022), and cells must produce approx. 6–10 new coccoliths between each cell division to ensure a complete cell covering for two daughter cells (based on mean C_N 18–20 in our samples). It would therefore take more than one generation of growth (> 24 h when $\mu < 0.7 \text{ d}^{-1}$) under new environmental conditions before C_L measurements start to reflect the increasing proportion of coccoliths of a different size produced in response to physiological adjustments to the new environment (e.g. salinity). However, coccoliths are produced quickly enough that coccolith size responses should be already evident after two generations, supporting the timescale

of morphological responses observed in our 156 h experiment. This might also explain why Iglesias-Rodríguez et al. (2017) did not observe any difference in C_L over a 72 h exposure of *E. huxleyi* strain NZEH to two different low-pH deep-seawater conditions, even though coccosphere volume was significantly different (+30 %) between the two deep seawater conditions after 72 h.

4.4 Intraspecific variability in *E. huxleyi* coccolith PIC – methodological considerations

Our estimates of coccolith PIC for strains PLYB11 (ranging from 0.04 to 0.38 pg C per coccolith) and RCC1232 (ranging from 0.06 to 0.39 pg C per coccolith) are comparable to the range of values reported for *E. huxleyi* morphotype A in the subantarctic zone of New Zealand by Saavedra-Pellitero et al. (2023; 0.02–0.35 pg C) and *E. huxleyi* in South Atlantic surface sediment samples (0.02–0.58 pg C; Horigome et al., 2014). However, our mean coccolith PIC values under each salinity treatment (Table 1) are at the lower end of the range of coccolith PIC values reported for *E. huxleyi* morphotype A coccoliths on the Patagonia shelf by Poulton et al. (2011; 0.12–0.42 pg C) and of morphotype A strains in culture (0.12–0.36 pg C under a range of temperatures; Rosas-Navarro et al., 2016, and 0.49–0.94 pg C in nutrient-replete cultures; Müller et al., 2012).

To assist with the comparison of our dataset and other published coccolith PIC estimates for *E. huxleyi*, we briefly address some of the factors that may contribute to species-specific variability in coccolith and coccosphere PIC estimates between publications. Firstly, we have used a well-established morphometric-based approach to estimate coccolith and coccosphere calcite (Young and Ziveri, 2000), which is a function of coccolith size and a species-specific shape factor. Any difference in coccolith size between cultures (e.g. due to strain-specific morphological differences or coccolith size responses to environmental treatments) therefore influences morphometric-based coccolith PIC estimates. For example, Southern Ocean *E. huxleyi* morphotype A isolates have larger mean coccolith size (3.19–3.48 μm ; Valença et al., 2024) compared to the mean coccolith size of the Norwegian and Mediterranean strains used here (2.54–3.24 μm ; Table 1), which contributes to the higher morphometric-based estimates of mean coccolith PIC for Southern Ocean strains (0.22–0.28 pg C; Valença et al., 2024) compared to PLYB11 (0.11 pg C) and RCC1232 (0.19 pg C). Here, we have also used previously determined strain-specific and salinity-condition-specific shape factors in our morphometric-based coccolith PIC estimates (Linge Johnsen et al., 2019) rather than a general *E. huxleyi* shape factor (for instance, $K_s = 0.02$ is recommended by Young and Ziveri, 2000). However, shape factors can vary within a species by up to 20 % (Young and Ziveri, 2000), and coccolith calcite mass can be decoupled from changes in coccolith size (Linge Johnsen and Bollmann, 2020a), factors that can

introduce uncertainty in morphometric-based coccolith PIC estimates. Coccolith PIC estimates from natural populations also span a greater range of genetic and phenotypic variability (and therefore coccolith size) than are measured in strain-specific laboratory cultures, which will also introduce variability in mean coccolith PIC for the same morphotype.

Morphometric-based estimates of coccolith PIC are generally in good agreement with estimates derived from other commonly applied methods, including birefringence-based thickness estimates of coccolith mass (Linge Johnsen and Bollmann, 2020b; Valença et al., 2024). Whilst a detailed discussion of alternative methods for estimating coccolith PIC is beyond the scope of our study, for the purpose of contrasting our estimated coccolith PIC values with published data derived from alternative methodologies, we note that coccolith PIC values derived from birefringence-based approaches have limitations related principally to coccolith crystallography and the choice of microscopy settings and calibration methods (e.g. Bollmann, 2014; Beaufort et al., 2014, 2021; González-Lemos et al., 2018; Linge Johnsen and Bollmann, 2020b). For instance, estimates of coccolith PIC derived from the birefringence-based method first published by Beaufort (2005) require a correction of the coccolith length because of the limited detection of the outer edge of the distal shield of *E. huxleyi* coccoliths when observed under cross-polarised light microscopy (e.g. Beaufort et al., 2008). However, the correction of the underestimated length by a factor of 1.25 (Beaufort et al., 2008; D’Amario et al., 2018) may subsequently lead to an overestimation of coccolith calcite content, as the undetected mass of the coccolith is estimated based on the birefringence of the thickest part of the coccolith. For further details, refer to Bollmann (2014) and Linge Johnsen and Bollmann (2020b). It has previously been noted that estimates of *E. huxleyi* coccolith calcite from birefringence-based methods can be higher than morphometric-based estimates of coccolith PIC made on the same morphotype and/or in similar regions (Poulton et al., 2011; Saavedra-Pellitero et al., 2023). Such methodological considerations contribute to intraspecific variability in coccolith PIC reported across different studies, in addition to the effect of any environmental treatment on morphology.

4.5 Implications of rapid responses to salinity stress

Whilst our morphological results represent changes over just 2–4 generations of growth under salinity stress, a general pattern of smaller cells with smaller coccoliths under hypersaline conditions (reduced cellular PIC with comparable cellular PIC : POC) and larger cells with larger coccoliths under hypersaline conditions (increased cellular PIC with comparable cellular PIC : POC) emerges from both strains investigated here (Fig. 6, Table 1). A similar pattern of smaller coccoliths under lower salinity and larger coccoliths under higher salinity was also found in acclimated cultures of several different *E. huxleyi* strains, including some open-ocean

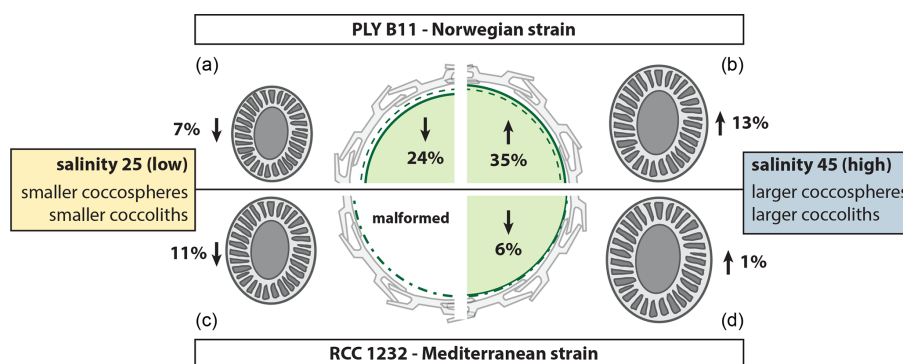


Figure 6. Summary of the morphological response of *E. huxleyi* strains PLYB11 (a, b) and RCC1232 (c, d) to abrupt exposure to low-salinity 25 conditions (a, c) and high-salinity 45 conditions (b, d) relative to coccosphere volume (centre cell illustration) and coccolith size (coccolith illustrations) under control salinity 35 conditions. Percentage change values are calculated based on mean C_L and mean \varnothing measurements at 148 h for PLYB11 and 140 h for RCC1232, which represent approx. 6 d growth under the salinity condition (see also Table 1). The dashed green line within the cell illustration indicates the equivalent cell size of mean cell volume under salinity 35 conditions at 148 or 140 h for PLYB11 and RCC1232, respectively.

strains (Green et al., 1998; Paasche et al., 1996; Fielding et al., 2009; Linge Johnsen et al., 2019; Saruwatari et al., 2016), in plankton samples (Bollmann et al., 2009), and in sediment core-top samples (Bollmann and Herrle, 2007).

The response of coccolith morphology and coccosphere (cell) size to the abrupt change in salinity conditions equates to a rapid (within hours to days) calcification response of *E. huxleyi* to salinity stress, with lower cellular PIC under hyposaline conditions, increased cellular PIC under hypersaline conditions, and similar cellular PIC:POC across all salinity conditions, due to the comparable timing of cell (coccosphere) size responses to salinity stress (Table 1). Our results show that *E. huxleyi* morphology and calcification is therefore sensitive to even relatively short-term (days to weeks) intervals of abrupt salinity change and, based on evidence from the literature, that these morphological effects are sustained when the new salinity condition persists for weeks to months (as would be typical in experiments using acclimated cultures).

Changes in coccolith size and coccolith morphology more generally are widely used as proxies for past environmental conditions through time (e.g. Bollmann, 1997; Henderiks and Bollmann, 2004), including paleosalinity (Bollmann et al., 2009; Herrle et al., 2018; Bollmann and Herrle, 2007). The response of *E. huxleyi* coccolith size to salinity stress within hours of exposure in our experiments signifies that transient and extreme salinity events will affect coccolith size in plankton samples alongside longer-term seasonal-to-decadal salinity trends. Seasonally, calcification responses on such short time frames may influence surface ocean alkalinity and inorganic carbon export in regions where *E. huxleyi* is a major component of the phytoplankton community. The analysis of coccolith size through time from marine sedimentary records remains most suited to capturing short- to longer-term salinity fluctuations, as sedimentation rates in

laminated sediments capture seasonal-to-annual timescales as well as sedimentation rates of thousands to hundreds of thousands of years per centimetre of sediment in deep-ocean sediments. However, we emphasise that the rapid morphological response of *E. huxleyi* to salinity (shown here) and other environmental variables, including exposure to CO_2 concentrations (Barcelos e Ramos et al., 2010) and distinct water masses (Iglesias-Rodriguez et al., 2017), already influence morphology and calcification at a cellular level within only one or two generations of growth. Relative to the timescale of interest, variability in the magnitude and timing of *E. huxleyi* calcification responses to salinity (and likely other environmental parameters) will therefore contribute to the natural within-species variability in morphology and biogeochemical traits observed in natural *E. huxleyi* populations.

5 Conclusions

The coccolithophore *E. huxleyi* has a naturally broad salinity tolerance, thriving in both relatively stable open ocean settings and the more variable environmental conditions of shelf seas and coastal regions. Despite this salinity tolerance, the physiology and morphology of *E. huxleyi* is responsive to changes in salinity. Our experiments show, for the first time, that measurable differences in *E. huxleyi* coccolith size and coccosphere size occur within hours of abrupt exposure to hypo- and hypersaline conditions. The resultant impact of these rapid morphological responses for cellular calcification on short timescales may impact surface ocean carbonate chemistry in regions where *E. huxleyi* is a dominant constituent of phytoplankton communities and contributes to the natural morphological variability in *E. huxleyi* coccoliths in the sedimentary record. The magnitude and timing of the response of *E. huxleyi* to salinity stress is strain-specific and may be related to different osmoregulation capacities

between strains, although further exploration of the physiological and biochemical mechanisms underpinning our results was beyond the scope of this study. Further insights would be gained from investigating the magnitude and timing of short-term morphological responses of a broader range of open-ocean and coastal *E. huxleyi* strains to a range of moderate-to-extreme salinity stress whilst measuring additional physiological indicators (e.g. photophysiology, carbon fixation rates, coccolith geochemistry, and cellular concentrations of DMSP and other solutes). Similarly, it is unclear whether other coccolithophore species respond to the onset of salinity stress in the same way, especially as the biogeography of many species is largely restricted to open-ocean settings with a smaller natural salinity range. If trends between morphology and salinity conditions are identified for modern representatives of longer-lived coccolithophore genera, the geological periods over which coccolith morphology can be used as an independent paleosalinity proxy could potentially be extended.

Data availability. Data reported in this study and Fig. S1 (SEM images of coccospheres) are available on Zenodo as Gebühr et al. (2024), <https://doi.org/10.5281/zenodo.11186906>. The dataset of carbonate system parameters at the start and end of experiments is available in the supplementary dataset (Table S4) of Gebühr et al. (2021), <https://doi.org/10.1371/journal.pone.0246745>.

Author contributions. The study was conceived by JOH, CG, and JB. CG planned and conducted the experiments and data collection. Morphometric data collection was performed by CG and RMS. Data analysis and interpretation were performed by RMS. The paper was written by RMS with contributions from all authors.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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