Abundances and morphotypes of the coccolithophore *Emiliania huxleyi* in southern Patagonia compared to neighbouring oceans and Northern Hemisphere fjords

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Abstract. Coccolithophores are potentially affected by ongoing ocean acidification, where rising CO2 lowers seawater pH and calcite saturation state (Ωcal). Southern Patagonian fjords and channels provide natural laboratories for studying these issues due to high variability in physical and chemical conditions. We surveyed coccolithophore assemblages in Patagonian fjords during late spring 2015 and early spring 2017. Surface Ωcal exhibited large variations driven mostly by freshwater inputs. High-Ωcal conditions (max. 3.6) occurred in the Archipelago Madre de Dios. Ωcal ranged from 2.0–2.6 in the western Strait of Magellan and 1.5–2.2 in the inner channel and was subsaturating (0.5) in Skyring Sound. *Emiliania huxleyi* was the only coccolithophore widely distributed in Patagonian fjords (> 96% of total coccolithophores), only disappearing in the Skyring Sound, a semi-closed mesohaline system. Correspondence analysis associated higher *E. huxleyi* biomasses with lower diatom biomasses. The highest *E. huxleyi* abundances in Patagonia were in the lower range of those reported in Norwegian fjords. Predominant morphotypes were distinct from those previously documented in nearby oceans but similar to those of Norwegian fjords. Moderately calcified forms of *E. huxleyi* A morphotype were uniformly distributed throughout Patagonia fjords. The exceptional R/hyper-calcified coccoliths, associated with low Ωcal values in Chilean and Peruvian coastal upwellings, were a minor component associated with high Ωcal levels in Patagonia. Outlying mean index (OMI) niche analysis suggested that pH and Ωcal conditions explained most variation in the realized niches of *E. huxleyi* morphotypes. The moderately calcified A morphotype exhibited the widest niche breadth (generalist), while the R/hyper-calcified morphotype exhibited a more restricted realized niche (specialist). Nevertheless, when considering an expanded sampling domain, including nearby southeast Pacific coastal and offshore waters, even the R/hyper-calcified morphotype exhibited a higher niche breadth than other closely phylogenetically related coccolithophore species. The occurrence of *E. huxleyi* in naturally low pH–Ωcal environments indicates that its ecological response is plastic and capable of adaptation.

1 Introduction

Coccolithophores are small unicellular phytoplankton (3–30 µm) bearing intricate plates called coccoliths formed of calcite, the more stable form of calcium carbonate (CaCO3) (Monteiro et al., 2016). Coccolithophores can carry out a
substantial portion of CaCO₃ precipitation in pelagic systems (Broecker and Clark, 2009), which may enhance oceanic matter export by CaCO₃ ballasting (Klaas and Archer, 2002), while also lowering alkalinity (Zondervan et al., 2001) and contributing to the carbonate counter pump (Passow and Carlson, 2012). Thus, understanding how coccolithophores respond to environmental stressors, such as ocean acidification (OA) due to rising pCO₂, is necessary for predicting future ocean biogeochemistry.

Chemically, calcite stability in seawater can be parameterized using its saturation state, defined as 
\[
\Omega_{\text{cal}} = \frac{[Ca^{2+}] [CO_3^{2-}]}{K_{sp,\text{cal}}}
\]
where \(K_{sp,\text{cal}}\) is the solubility constant for calcite), which is decreased by OA. When \(\Omega_{\text{cal}} < 1\), calcite dissolution is thermodynamically favoured, whereas calcite precipitation is favoured when \(\Omega_{\text{cal}} > 1\). Most of the ocean surface is predicted to remain supersaturated with respect to calcite in the future ocean (Feely et al., 2009), although some coastal zones may experience \(\Omega_{\text{cal}} < 1\) in the euphotic zone either due to increased pCO₂ in areas of naturally high CO₂ upwelling (e.g., Franco et al., 2018) or due to freshening (e.g., Tynan et al., 2014). Furthermore, drops in \(\Omega_{\text{cal}}\) may negatively affect biological calcite production rates even before becoming undersaturated (Doney et al., 2009).

In contrast to other calcifying organisms, coccolith formation occurs intracellularly in Golgi-derived vesicles, involving sustained fluxes of Ca²⁺ and dissolved inorganic carbon (primarily HCO₃⁻) from the external medium and high H⁺ efflux to maintain cellular pH homeostasis (reviewed by Taylor et al., 2017). Which extracellular carbonate chemistry parameter most influences intracellular coccolithophore calcification is debated, e.g., whether \(\Omega_{\text{cal}}\) or more complex relationships involving HCO₃⁻, H⁺, and CO₂ (Bach et al., 2015; Cyronak et al., 2016; Kottmeier et al., 2016; Gafar et al., 2018). Additionally, OA can have contrasting effects, with increased CO₂ availability potentially benefiting photosynthesis but high H⁺ negatively affecting metabolisms besides calcification (Kottmeier et al., 2016; Paul and Bach, 2020). A wide range of calcification and growth responses to OA have been reported in laboratory studies of coccolithophores, mostly using the cosmopolitan and most abundant species *Emiliania huxleyi* (reviewed in Meyer and Riebesell, 2015). With some notable exceptions (e.g., Iglesias-Rodriguez et al., 2008), most culture studies showed reduced calcification rates of *E. huxleyi* in response to simulated OA, while there is no clear trend on growth rates. In a mesocosm experiment using a Norwegian fjord community, increased pCO₂ levels (> 500 μatm) resulted in lower growth rates of *E. huxleyi*, preventing it from blooming (Riebesell et al., 2017). This result contrasted with a long-term observational study showing a steady increase in coccolithophore stocks (related to diatoms and dinoflagellates) from pre-industrial to present-day high pCO₂ levels in the North Atlantic (Rivero-Calle et al., 2015), although both studies predict a decrease in net calcification.

Morphological variability in *E. huxleyi* has been reported with several morphotypes described so far with different degrees of calcification of the coccoliths, such as fusion of coccolith elements or calcite overgrowth (Young et al., 2003). Morphotype-A coccoliths have a grill central area and tend to be moderately calcified, while morphotypes B and C have more lightly calcified distal shield elements, and the central area is either a plate or open (type O) (Young and Westbroek, 1991; Hagino et al., 2011). Additional morphotypes, or morphotype sub-classes, include B/C (intermediate in coccolith size between B and C) and R (*Reticulofenestra*-like), considered an A morphotype where distal shield elements are mostly or completely fused (Hagino et al., 2011). However, the readily identifiable qualitative distinctions may not easily translate into quantitative differences in the calcite produced per cell in the ecosystem, as for example, due to variability in the rate of coccoliths produced per cell (Johnsen and Bollmann, 2020).

Nevertheless, cultured isolates maintain their morphotype classifications even under variable environmental conditions that can alter total calcite production and even lead to coccolith malformation (e.g., Young and Westbroek, 1991; Langer et al., 2011; Müller et al., 2015; von Dassow et al., 2018; Mella-Flores et al., 2018), suggesting a genetic determination of coccolith morphology. One genetic marker has been associated with morphological variability in *E. huxleyi*. The calcium-binding protein GPA has been potentially associated with *E. huxleyi* coccolith deposition (Corstjens et al., 1998). Although the function of this protein is unclear, the 3′ untranslated region (non-coding) showed consistent differences between morphotypes, with all morphotypes A and R showing alleles (coccolith morphology motifs) CMM I, III, or IV and B, C, and C morphotypes showing CMM II (Schroeder et al., 2005; Krueger-Hadfield et al., 2014). The uronic acid content of coccolith-associated polysaccharides also varies among strains, and the one R morphotype tested was much higher in this character than most of the other A morphotypes (Rickaby et al., 2016). It is likely that further comparative biochemical analyses following Rickaby et al. (2016) and/or associating comparative genomics analyses (e.g., studies such as Read et al., 2013; von Dassow et al., 2015; Bendif et al., 2019) with morphometric analyses may identify genetic markers associated with sub-types within the broader A and B morphotypes. However, mitochondrial phylogenies classify *E. huxleyi* into a warmer water clade and a colder water clade, and each clade contains both A (including R) and B (or B/C or O) morphotypes (Hagino et al., 2011; Bendif et al., 2014), and B/C morphotypes also occurred in different genetic groups defined by microsatellite markers (Krueger-Hadfield et al., 2014), although another microsatellite study did find a separation between A and B/C morphotypes (Cook et al., 2013). Therefore, different morphologies likely correspond to stable genetically determined phenotypes that might reflect adaptations selected to specific
conditions within a taxon whose recent evolution has been as a single biological species (Filatov, 2019).

In a global survey, Beaufort et al. (2011) found a general pattern of decreasing calcification with increasing pCO$_2$ and a concomitant decrease in CO$_2$. Interestingly, calcite mass variability was predominately the result of assemblage shifts both among closely related species and among morphotypes of the same species, from predominance of large and heavily calcified *Gephyrocapsa oceanica* cells, through intermediate moderately calcified *E. huxleyi* (A morphotype), to more lightly calcified *E. huxleyi* cells (B/C or C morphotype). In the subtropical and tropical eastern South Pacific, an exceptionally hyper-calcified R morphotype of *E. huxleyi* (henceforward referred to as R/hyper-calcified, showing the complete fusion of distal shield elements and partial or complete calcite overgrowth of the coccolith central area) was dominant in coastal upwelling waters with relatively high pCO$_2$ and low $\Omega_{\text{calc}}$ levels, causing an inversion in the trend of calcite-$\text{per-coccolith vs. } \Omega_{\text{calc}}$ seen in the rest of the ocean (Beaufort et al., 2011; von Dassow et al., 2018). Likewise, Smith et al. (2012) observed an increase in the proportion of *E. huxleyi*, corresponding to an “overcalcified” morphotype (with complete overgrowth of the coccolith central area but without fusion of distal shield elements, referred to hereon as A-CC for covered central area) that occurred during the winter decline of $\Omega_{\text{calc}}$ in the Bay of Biscay (North Atlantic). These results suggested that perhaps the A-CC and R/hyper-calcified morphotypes are likely resistant to low $\Omega_{\text{calc}}$. However, while the B/C morphotype was reported to be especially sensitive to low $\Omega_{\text{calc}}$ compared to moderately calcified and overcalcified strains of morphotype A (Müller et al., 2015), other experimental results did not find a higher resistance of the R/hyper-calcified subtype to high CO$_2$ and low $\Omega_{\text{calc}}$ when compared to moderately calcified strains of morphotype A isolated from neighbouring waters (von Dassow et al., 2018). These results highlight that it is unclear how the physiological effects of OA on coccolithophores in culturing conditions translate to community-level responses in the field.

Fjord systems provide especially interesting natural laboratories for observing how coccolithophores may be affected by environmental conditions due to high spatial and seasonal variability in chemical and biotic conditions. In the Norwegian fjord system and the neighbouring North Sea, *E. huxleyi* has been very well studied for over a century, where it forms dense spring blooms but is also prominent throughout the year (Birkenes and Braarud, 1952; Berge, 1962; Kristiansen et al., 1994; Fernandez et al., 1996; Egge et al., 2015). Southern Patagonia, on the Pacific side of South America, hosts the largest network of fjords and channels in the world. Aquatic ecosystems of southwest Patagonia (50–55°S) are dominated by the transition between oceanic and estuarine–brackish waters. Denser, saltier, nitrate- and phosphate-rich but silicate-poor Sub-Antarctic Surface Water (SAASW) intrudes below nitrate- and phosphate-poor but silicic-acid-rich surface waters influenced by substantial freshwater inputs (copious precipitation, rivers, glacier melt; Dávila et al., 2002; Sievers and Silva, 2008; Torres et al., 2014). Generally, surface waters with low salinity and low alkalinity are undersaturated in dissolved CO$_2$ during the spring and summer seasons (Torres et al., 2011). The Archipelago Madre de Dios (AMD) is an interesting exception to this pattern, where extreme precipitation/runoff in the “limestone” basin on the western AMD produces surface waters with low salinity and high alkalinity while maintaining low dissolved silicate compared with the eastern basin (Torres et al., 2020). These gradients create a unique contrast for exploring the influence of chemical conditions on the ecology of calcified phytoplankton, as changes in $\Omega_{\text{calc}}$ are driven mainly by freshening rather than upwelling of high pCO$_2$.

In contrast to the Norwegian fjord system, *E. huxleyi* blooms have not been reported in Patagonian fjords; however information on coccolithophores in these waters is scarce. A study documenting coccolithophores in the Strait of Magellan found that this group represented a minor fraction of the small-sized phytoplankton (Zingone et al., 2011), but other published studies have not specifically sampled for coccolithophores. The Patagonian shelf on the Atlantic side experiences large *E. huxleyi* blooms (Poulton et al., 2011, 2013), but satellite observations suggest coccolithophore blooms are of lower intensity in the Pacific waters to the west (Hopkins et al., 2019). These observations raise the question of how coccolithophore communities on the western coast and fjords–channels of Patagonia compare with nearby oceans and with fjord systems in the Northern Hemisphere. Here, we evaluated how physical, chemical, and biological features influence the distribution, abundance, and biomass of coccolithophores as well as the proportions of *E. huxleyi* morphotypes of varying calcification levels throughout southern Patagonia fjords. In particular, three research questions were addressed. (i) What coccolithophore assemblages and *E. huxleyi* morphotypes are present in fjords and channels of southern Patagonia? (ii) How do these morphotypes and the co-occurring phytoplankton (mostly diatoms) vary with physical and chemical factors? (iii) Do the abundance and relative composition of *E. huxleyi* morphotypes reflect populations in adjacent Pacific, Atlantic, or Southern Ocean waters or instead exhibit similarities to the Norwegian fjord system?

2 Materials and methods

2.1 Sampling

Two cruises were conducted on board the vessel M/N *Forrest* during the late austral spring 2015 (26 November to 3 December) and early austral spring 2017 (22 to 28 September) in southern Chilean Patagonia (∼50–54°S, 71–76°W). The sampling track of 2015 was from the Archipelago Madre de Dios (AMD), crossing the inner channel (IC) to the western part of the Strait of Magellan (WSM), and entering eastward
Figure 1. Maps of southern Patagonia showing the study sites and stations sampled during the austral late spring 2015 (a) and early spring 2017 (b). Salinity recorded at the surface during the two cruises is plotted. The approximate perimeter of the Southern Ice Fields (SIF) is depicted. A zoom into the Archipelago Madre de Dios (AMD) zone with salinity and CaCO$_3$ surface values recorded in 2015 is provided in Fig. S1. Maps were produced by Ocean Data View (Schlitzer, 2018).

into the Otway and Skyring sounds (OS, SS) to end up near Punta Arenas (Fig. 1a). The 2017 sampling was from the interior WSM, crossing the IC, and ending in the AMD zone (Fig. 1b). In both cruises, surface water (< 5 m) was pumped continuously on board every 15–20 min for determination of salinity and temperature with a YSI-30 termosalinometer (Yellow Springs, OH, USA) and pCO$_2$ with a Qubit-S157 CO$_2$ analyzer (Kingston, Ontario, Canada). The CO$_2$ analyzer was calibrated daily with 0 ppm CO$_2$ (air treated with soda lime) and 403 ppm air–CO$_2$ mixture standard (Indura, Chile). Surface samples for determination of the planktonic assemblages and chemical variables (i.e., concentration of macronutrients, opal, total chlorophyll $a$, and the carbonate system parameters) were collected at discrete sampling stations (Fig. 1a–b). Twenty-one stations were sampled in 2015: five in the AMD (Sts. 1–5), four in the IC (Sts. 6–9), nine in the WSM (Sts. 10–12, 16–21), two in the OS (Sts. 13 and 14), and one in the SS (St. 15). Eleven stations were sampled in 2017: three in the AMD (Sts. 30–32), five in the IC (Sts. 25–29), and three in the WSM (Sts. 22–24).

Conductivity, temperature, and depth (CTD) vertical profiles were additionally obtained at selected localities on both cruises. In 2015, three casts were performed into the AMD zone and one cast into the SS using a CTD Seabird 19 plus (Sea-Bird Scientific, Bellevue, WA, USA) equipped with photosynthetically available radiation (PAR) and oxygen sensors. Two profiles were performed in 2017 in the AMD zone, by the deployment of a CTD Seabird 25 plus with PAR and oxygen sensors. The depth of 1% of penetration of PAR (euphotic zone) was calculated from maximum surface PAR values. CTD profile binning was 1 m. In both years, samples for the determination of planktonic assemblages and chemical variables were obtained at discrete depths using 5 L Niskin bottles to which the CTD profiles were attached (depths pre-determined from prior studies in the region, aiming to adequately sample the surface mixed layer, pycnocline, and vertical variation in chlorophyll fluorescence). Complete environmental and biological data are provided in the Supplement (Tables S1–S4).

2.2 Plankton assemblages

Samples for the determination of planktonic organisms through the Utermöhl (1958) method were collected only in 2015. For that, duplicate 100 mL water samples were pre-filtered through 200 µm Nitex mesh, fixed with a formaldehyde–glutaraldehyde solution (1% formaldehyde, 0.05% glutaraldehyde, 10 mM borate pH 8.5) and stored at 4°C. In the laboratory, water samples were brought to room temperature, gently homogenized, and sedimented into 100 mL chambers for 24–48 h before counting and identification. The absolute abundances of the microplankton (20–200 µm in size) and coccolithophores (~6 µm in diameter) were estimated with an inverted microscope (Olympus CKX41) connected to a digital camera (Motic 5.0). For counts of diatoms, dinoflagellates, and other planktonic cells greater than about 40 µm, the whole chamber was examined...
of water analyzed. The coccolithophore abundances were and E. huxleyi at room temperature. For the identification of coccolithophores 0.8 µm polycarbonate filters that were subsequently dried at 300 mL of surface water, immediately after sampling, onto microscopy (SEM) analysis were obtained by filtering 200–3.5 mm 1500 magnification. Counts of total coccolithophores were performed with a 40× objective with cross-polarized light (Edmund Optics polarizers 54 926 and 53 347).

In both cruises, samples for the identification and quantification of coccolithophores through scanning electron microscopy (SEM) analysis were obtained by filtering 200–300 mL of surface water, immediately after sampling, onto 0.8 µm polycarbonate filters that were subsequently dried at room temperature. For the identification of coccolithophores and E. huxleyi morphotypes, a portion of each dried filter was cut, sputter-coated with gold, and examined in either a Quanta FEI 250 or Quanta FEG 250 SEM (both FEI, Hillsboro, Oregon, USA). As water samples for light microscopy counts were not available for two samples from 2015 (St. 3 and St. 5 at 8 m) and all samples from 2017, total coccolithophore abundances were obtained from SEM counts for those samples. On average, 70 images per filter were captured at 1500× magnification (276 × 184 µm per frame), covering 3.5 mm² of the filter area corresponding to 1.8–3.4 mL of water analyzed. The coccolithophore abundances were calculated using the following equation:

\[
cells \, L^{-1} = \frac{\text{effective filtration area (mm²) × total number of counted cells}}{\text{analyzed filtered area (mm²) × volume of filtered water (L)}}
\]

To check for differences in coccolithophore counts obtained through sedimentation and inverted light-microscopy versus filtration and SEM examination (hereinafter SEM and Utermöhl counts, respectively), polycarbonate filters from three selected Utermöhl samples (showing higher, medium, and lower coccolithophore abundances) were analyzed with SEM as outlined above. Coccolithophore SEM counts were consistently about twice as high compared to Utermöhl counts (average 1.84), agreeing with the correction factor suggested by Bollmann et al. (2002). Thus, all total coccolithophore counts obtained by the Utermöhl method were multiplied by 1.84 to be comparable to SEM counts. To estimate the absolute abundances at species and morphotype level, the relative abundance of each coccolithophore species or E. huxleyi morphotype determined from SEM counts was multiplied by the absolute abundance of total coccolithophore cells. Saturation curves confirmed that the number of analyzed coccospheres (minimum 40 coccospheres per sample) was enough to capture the specific/morphotype diversity.

SEM images taken at 20 000–25 000× magnification were used to categorize E. huxleyi cells in the different morphotypes according to the morphology of distal shield and the central plate of the coccoliths (following Young and Westbroek, 1991; Young et al., 2003; Hagino et al., 2011; von Dassow et al., 2018). Given high morphological similarities in the A morphotype coccoliths with those found by Young (1994) in Norwegian fjords, they were here classified as lightly, moderately, and robustly calcified, based on the morphology of distal shields and central plates (Fig. 2; Table 1). Moreover, two extremely heavily calcified A morphotypes were observed: the A-CC (with closed central area but distal shield elements mostly not fused) and the R/hyper-calcified forms (bottom) are shown. For statistical analysis, the moderately and robustly calcified A morphotypes were merged under “moderately calcified A morphotype”, and the few O and C specimens were categorized into the lightly calcified subgroup. Scale bar = 1 µm.

![Figure 2](image-url)
Table 1. Classification of *E. huxleyi* coccospheres based on the calcification level of coccoliths. Mean ± standard deviations of distal shield length and coccosphere diameter of morphotypes observed in Patagonia are given for coccospheres where at least one distal shield was visible and adequately oriented (numbers in parentheses).

| Morph | Calcification | Distal shield and central area distinguishing features | Distal shield length (µm) | Coccosphere size (µm) | Comparable morphotype | Reference |
|-------|---------------|-----------------------------------------------------|---------------------------|-----------------------|-----------------------|-----------|
| A     | Light         | Delicate and well separated distal shield elements (<50% of distal shield elements are slits), which end up in a wide central area consisting of plate or lath-like elements | 3.3 ± 0.3 (15) | 5.7 ± 0.5 (15) | Under-calculated | Young (1994) |
| A     | Moderate      | Straight low-profile distal shield formed by thicker elements (<25% of distal shield area is open between elements) fused to tube elements delimiting a grilled cleanly to semi-open central area | 3.5 ± 0.3 (430) | 6.3 ± 0.9 (430) | Typical A morphotype | Young and Westbrook (1991) |
| A     | Robust        | Robust calcification of elements (<25% of distal shield area is open between elements) from partially to nearly completely fused extending from the outer rim to a wide to narrow central area delimited by robust tubes | 3.3 ± 0.3 (259) | 5.9 ± 0.7 (259) | A overcalculated Type A | Young (1994) |
| A     | CC            | Thicker to robust but not fused elements and central area completely or nearly completely covered | 3.0 ± 0.2 (26) | 5.7 ± 0.6 (26) | Overcalculated Type A | Henderiks et al. (2012) |
| A     | R/hyper       | Heavily calcified distal shield elements completely fused and central area grilled but partly or completely covered | 3.7 ± 0.4 (13) | 7.2 ± 1.0 (13) | Overcalculated R-type R overcalculated | Cubillos et al. (2007) |
| A     | Light         | Distal shield elements in low profile and greater coccosphere size compared with light A morphotype | 3.4 ± 0.2 (7) | 6.9 ± 1.4 (8) | Type B, Type B/C, Type C | Young and Westbrook (1991) |

* The Plate B–c of Saavedra-Pellitero et al. (2019) are grouped together as “overcalculated” in that reference but are distinguished in the present study as R/hyper-calculated and A-CC morphotypes. Plate 1e of Saavedra-Pellitero et al. (2019) corresponds to overcalculated A morphotype in Young (1994) and the present study. Similarly, the “Type A overcalculated” in Fig. 3c of Cubillos et al. (2007) corresponds to the A-CC morphotype here (as distal shield elements are not fused or only partly fused in most coccoliths) while Fig. 3d of the same reference, identified also as “Type A overcalculated” appears to show both nearly complete fusion of the distal shield element and nearly complete overcalcification covering the central area, thus corresponding to the R/hyper-calculated morphotype in the present study. B and C morphotypes are distinguished by distal shield diameters >4.5 or <3.5 µm, respectively, with B/C being intermediate. O variants have varied distal shield diameters with an empty (open) central area (Hagino et al., 2011). B and B/C were not observed in Patagonia. The lightly calcified *E. huxleyi* in the dataset of von Dassow et al. (2018) included a continuum of characteristics from A to B and B/C or C.

Although these malformed and collapsed coccoliths were observed in <9% of the morphotype-A cells, it was almost always possible to classify those abnormal coccospheres into one of the abovementioned morphotypes (Fig. S2). SEM images were also used to measure the orthogonal coccosphere diameters and, when available, coccolith distal shield length (ImageJ software version 1.4.8 for Mac OS).

Biovolumes (µm³) of *E. huxleyi*, diatoms, dinoflagellates, and naked flagellates were estimated assuming recommended geometric shapes (Hillebrand et al., 1999; see Table S4). For *E. huxleyi*, a spherical geometric shape was assumed, and the average of the minimum and maximum diameters was used for biovolume calculations. Biovolume calculations were then converted to carbon biomass by using regression equations proposed by Menden-Deuer and Lessard (2000) for diatoms (pg C cell⁻¹ = 0.288 x volume⁻⁰⁸¹¹) and other protists (pg C cell⁻¹ = 0.216 x volume⁻⁰⁹³⁹). We assumed a constant cytoplasm diameter to be 60% of the mean *E. huxleyi* coccosphere diameter (O’Brien et al., 2013), whereas cytoplasm volumes of 50% and 78% were used for diatoms and dinoflagellates, respectively (i.e., total cellular volume minus frustule or theca and vacuole volumes; Sicko-Goad et al., 1984). Absolute abundance data were standardized to cells per litre and multiplied by specific carbon contents per cell (pg cell⁻¹) to derive total carbon biomass (Total C, µg C L⁻¹). We used the biogenic silica concentration (µmol opal L⁻¹) as a proxy of diatom biomass along the 2017 track, as samples for microscopy counts were
not available. For this, the bSi concentration was converted into carbon units using the average net silicate-to-carbon ratio of 0.52 (mol/mol) found by Brzezinski et al. (2003) in the Southern Ocean. There was a significant linear relationship between diatom carbon biomass estimated with microscopy/allometry and that calculated from bSi concentration in 2015 samples ($R^2 = 0.60$, $p < 0.05$, slope $= 0.8$; $N = 11$), with an offset (16 µg CL$^{-1}$) likely from other contributors to bSi (e.g., silicoflagellates, radiolarians) as well as the contribution of Minidiscus spp. (data not shown) not included in microscopy/allometric carbon estimation. The presence or absence of diatoms was confirmed qualitatively in 2017 by SEM images at 1500× magnification, and a semi-quantitative evaluation was made as follows: low (few valves), intermediate (at least one species with several valves or chains), and high (many species with several cells or chains). It should be kept in mind that there can be substantial variation in diatom carbon biomass estimated by microscopy vs. bSi, due to variability in diatom C : Si ratios (Leblanc et al., 2018).

2.3 Chemical analyses

Macronutrients, opal, total chlorophyll $a$ (chl $a$), pH, and total alkalinity ($A_T$) were determined as described in Torres et al. (2020). Full carbonate system parameters (including $\Omega_{calc}$) were estimated from pH, $A_T$, salinity, temperature (25°C as input and in situ temperature as output conditions), and pressure (0 dbar as input and depth as output conditions) using CO2Sys Excel macro spreadsheet version 2.1 (Pierrot et al., 2006) with Mehrbach set of solubility constants (Mehrbach et al., 1973) refitted by Dickson and Millero (1987). To extrapolate full carbonate parameters from $pCO_2$ (onboard sensor) and salinity measurements where alkalinity samples were not directly available (due to mismatch in chemical and biological sampling along the IC-WSM 2015 track), the regression curve for the salinity–$A_T$ relationship ($\mu$mol kg$^{-1}$) = 63.4 × salinity + 101 ($R^2 = 0.99$; $N = 186$; Torres et al., 2020) was used to derive $A_T$ estimated from salinity. It is important to note that this relationship has been stable for over a decade in Patagonia (Torres et al., 2011, 2020). $pCO_2$ values delivered by the onboard sensor (underway sampling) correlated with $pCO_2$ calculated from $A_T$–pH pairs (discrete sampling) in the same 2015 samples ($R^2 = 0.56$, $p < 0.001$; $N = 17$), with an overestimation of 6 µatm (2%). The differences between measured and calculated $pCO_2$ values are small compared to the high ranges in the variability of $pCO_2$, salinity, $A_T$, and pH and should not affect the objectives of the present study. Exceptionally, the calculated $pCO_2$ values for SS were overestimated by up to 36% concerning $pCO_2$ measurements (comparing 15 readings from the sensor with three calculated values). This disagreement could be due to various local factors that increase the sensitivity of calculated $pCO_2$ to $A_T$ (Abril et al., 2015) or pH uncertainties due to differences in salinity between buffers and samples. Therefore, it should be kept in mind that in the case of SS the uncertainties in the carbonate system could be more substantial. Finally, in order to calculate the ratio (in mol/µmol) of bicarbonate ($HCO_3^-$) to proton ($H^+$), the $HCO_3^-$ was divided by the antilog$_{10}$ of pH (total $H^+$ scale).

2.4 Statistical analysis

All statistical analyses were performed in R software using packages freely available on the CRAN repository (R Core Team, 2019). As measurements for nitrate, phosphate, DSi, bSi, and chl $a$ in 2015 were scattered and uncoupled from plankton sampling, these variables were used only descriptively and not included in the statistical analyses.

2.4.1 Environmental gradients

Physical–chemical data obtained from the surface at the discrete sample stations in both 2015 and 2017 were combined in a unique matrix and standardized by subtracting the mean and dividing by the standard deviation (Legendre and Legendre, 2012). We used the varclus function in the “Hmisc” package based on Spearman’s correlation to detect redundant environmental variables ($N = 32$). Temperature, salinity, pH, and $\Omega_{calc}$ were selected as they are non-redundant based on Spearman’s correlation < 0.75 (Fig. S3), and they are easiest to interpret from a biological or cell physiological point of view. We also included CO$_2$. It was moderately correlated with pH (Spearman correlation = 0.8) but represents the substrate for photosynthesis and is typically incorporated as a driving variable in ocean acidification studies. HCO$_3^-$ may more directly impact sensitivity of coccolithophores in lab measurements (e.g., Kottmeier et al., 2016; Gafar et al., 2018), but it was redundant with the other variables. The selected standardized variables were then used in two separate cluster analyses to recognize groups of sampling stations with similar characteristics in 2015 ($N = 21$) and 2017 ($N = 11$). For that, Euclidean distance matrices were first calculated based on selected standardized variables ( vegdist function in the “vegan” package) and then included in hierarchical cluster analyses based on the Ward method using the hclust function in the basic “stats” package.

2.4.2 Testing for a relationship of Emiliania huxleyi vs. diatoms

To characterize the diatom species associated with the different E. huxleyi morphotypes and other coccolithophore species, we performed a non-metric multidimensional scaling (nMDS) using the metaMDS function in the vegan package, based on the log-transformed [ln ($x + 1$)] coccolithophore and diatom abundances (cells L$^{-1}$) in Patagonia (only 2015) and the other coastal and oceanic locations (von Dassow et al., 2018) ($N = 52$). The function heatmap of the basic stats package was then used to plot the abundance of
the coccolithophore and diatom species related to the clusters based on the nMDS scores of species/morphotypes and samples. As both the nMDS and OMI (see below) analyses suggested a clear separation between Patagonia fjords and the other coastal/oceanic areas, we used the IndVal analysis (Dufrène and Legendre, 1997) to identify indicator species for both areas, based on log-transformed abundances (indval function in “labdsv” package).

We aimed to assess how *E. huxleyi* and diatom biomasses were related to each other and with the environmental conditions throughout Patagonia fjords. However, the different methods used to estimate diatom biomass in both years precluded the use of *E. huxleyi*-diatom ratios. Moreover, the use of regression-based analyses was not recommended due to the absence of a linear relationship between *E. huxleyi* and the different physical–chemical variables. To overcome these limitations, we created three categorical variables for both *E. huxleyi* and diatom biomasses (low, intermediate, and high) based on their 25th, 25–75th, and > 75th percentiles, respectively. We then performed a correspondence analysis (CA) using the function cca in the vegan package, based on the presence or absence of these new categorical variables in each sample (*N* = 32), followed by fitting the standardized physical–chemical variables to the CA plot using the envfit function (10 000 permutations).

### 2.4.3 Niche analysis

We used the outlying mean index (OMI) analysis (Dolédec et al., 2000) to assess how the different physical–chemical variables (selected in Sect. 2.4.1) were associated with the realized niche of the different *E. huxleyi* morphotypes. The OMI index represents the marginality (i.e., niche position) and measures the distance between the average habitat conditions used by a given population and the average environmental conditions across the study area (represented by the point where the two multivariate axes intersect at zero). The tolerance (Tol) accounts for the dispersion of samples containing organisms of the population from the average environmental condition (i.e., niche breadth), whereas the residual tolerance (RToI) accounts for the proportion of the variability unexplained by the variables included in the analysis (Dolédec et al., 2000). Thereby, a species having a low OMI (species score close to zero, located in the centre of the multivariate space) and high Tol is likely one that utilizes a wider array of resources and maintains populations within a wider variety of conditions (i.e., generalist), when compared with the specialized and less resilient species with a more restricted realized niche associated with high OMI and low Tol that are expected to show lower Tol and may also be associated with lower OMI (Dolédec et al., 2000).

The OMI analysis was performed using the niche function in the “ade4” package (Dray and Dufour, 2007), simultaneously considering the data obtained for 2015 and 2017 (*N* = 32). To compare the patterns observed in Patagonia to other localities in the southeastern Pacific, a complementary OMI analysis was performed, including records of coccolithophore assemblages and *E. huxleyi* morphotypes from nearby coastal and oceanic waters (published by von Dassow et al., 2018) in addition to the data used in the first analysis (*N* = 64). A 1.84× correction factor (determined as informed in Sect. 2.2) was applied to these data, as coccolithophore counts from von Dassow et al. (2018) were obtained by the Utermöhl method. In both cases, data were arranged in two matrices, one containing the coccolithophore abundances and a second matrix with the standardized physical–chemical variables. Coccolithophore abundances were previously Hellinger-transformed (Legendre and Legendre, 2012). Since Hellinger transformation is obtained by the squared root of relative abundances, the potential biases from comparing data from both SEM and Utermöhl counts were minimized. The statistical significance of the morphotypes/species marginality was tested using the Monte Carlo method included in the ade4 package (10 000 permutations). The envfit function in the vegan package (Okansen et al., 2007) was then used to fit the five environmental variables to the OMI scores (10 000 permutations).

### 3 Results

#### 3.1 Late spring southern Patagonia 2015

The hierarchical clustering based on the surface values of the selected physical–chemical variables in the late austral spring 2015 (i.e., salinity, temperature, *Ω*<sub>cal</sub>, pH, and *p*CO<sub>2</sub>) showed a clear separation between the sampling station at the Skrying Sound (SS; St. 15) and the other localities (Fig. 3a). The other stations were grouped in two main clusters: one cluster composed of stations in the Archipelago Madre de Dios (AMD) and the inner channel (IC) and a second one composed mainly of stations in the western part of the Strait of Magellan (WSM). Samples from the Otway Sound (OS; Sts. 13–14) were distributed between the two clusters. The cluster separation seemed to be mainly related to temperature and salinity dissimilarities, while stations 4 and 15 differed from others by their relatively low *p*CO<sub>2</sub>–high *Ω*<sub>cal</sub> and high *p*CO<sub>2</sub>–low *Ω*<sub>cal</sub> conditions, respectively (Fig. 3b). Surface salinity ranged from > 29 in the AMD and southernmost WSM stations to as low as 17 in the SS (St. 15), with intermediate values throughout the IC and in the OS (range: 26–29; Fig. 3a–b). A north–south gradient of decreasing surface temperature was recorded from 9.0–10.0 °C around the AMD zone to 7.1 °C near Helado Sound (Sts. 17–18; Figs. 3a–b, S4a). Surface waters were mostly undersaturated relative to atmospheric *p*CO<sub>2</sub> (< 400 µatm) with minimum values at St. 4 in the AMD (241 µatm). *CO*<sub>2</sub> oversaturation was only observed at the SS (542 µatm). Similar to salinity, *Ω*<sub>cal</sub> varied widely, ranging from highly saturated conditions in the AMD zone (*Ω*<sub>cal</sub> range: 2.5–3.6, pH...
in situ range: 8.03–8.21), intermediate levels in the interior WSM (Ω$_{cal}$ range: 2.4–2.6; pH range: 8.04–8.07), lower levels in the southern IC zone (Ω$_{cal}$ 2.0–2.2, pH ~ 8.0), and a subsaturated extreme reached at the SS sampled site (Ω$_{cal}$ 0.5, pH 7.73, Figs. 3a–b, S5a). Moderate to low DSi (< 6 µm) and nitrate (NO$_{-3}$ < 6 µm) were recorded in southern Patagonia, dropping to a minimum in the southern IC zone (Table 2, Fig. S6).

The AMD and IC zones showed relatively low phytoplankton biomasses (< 20 µg C L$^{-1}$) dominated by naked flagellates (Fig. 3c–d), whereas diatoms were associated with higher phytoplankton biomasses (> 40 µg C L$^{-1}$) in Eleuterio Channel (St. 4; mostly chains of Chaetoceros spp.) and the WSM (Sts. 12, 16–21; mostly chains of Leptocylindrus spp., Chaetoceros spp., and Thalassiosira spp.). The contribution of dinoflagellates to biomass was highest in IC St. 8, OS Sts. 13–14, and WSM Sts. 12 and 16. Coccolithophores only accounted for 0.2%–12.8% of total C biomass (0.1–4.0 µg C L$^{-1}$), reaching > 6% only in the southern IC (Sts. 8–10) where diatom biomasses were among the lowest observed. *E. huxleyi* dominated the coccolithophore assemblages in all samples (> 98%), with a few Syracosphaera spp. coccospheres (mostly collapsed) found at the AMD (Table S2). *E. huxleyi* abundances (Fig. 3e) ranged from 0 to 2.76 × 10$^5$ cells L$^{-1}$, being most abundant in southern IC (> 1.03 × 10$^5$ cells L$^{-1}$ in Sts. 7–10) and only absent from SS (St. 15), which was the only station where conditions were undersaturated with respect to calcite (Ω$_{cal}$ < 1). *Emiliania huxleyi* populations were mostly composed of the moderate A morphotype (which also included cells of the robustly calcified A morphotype; see Methods) (Fig. 3e, Table 3). Few lightly calcified A cells were observed. Among all samples, only eight total O and C and no B or B/C morphotype cells were detected. The heavily A-CC and R/hyper-calcedified morphotypes were restricted to the AMD zone.

Three vertical profiles were performed in the AMD estuarine zone (Fig. 4): one at the western limestone AMD basin (St. 3), one between the western and eastern AMD basins (St. 4), and one at the easternmost basin (St. 5, Figs. 4, S1). All samples were taken within the euphotic zone (1 % PAR), which extended down to 36 and 30 m at Sts. 3 and 4, respec-
Table 2. Physical and chemical conditions and photosynthetic and silicified biomass proxies in the surface waters (<5 m) of southern Patagonia fjords. Mean, standard deviation (SD), and range of each variable and number of samples for late spring 2015 and early spring 2017 are shown. Only values matching planktonic samples discussed in the text are included, except for chlorophyll a (chl a), opal, nitrate, and silicate 2015 for which values are between 3–28 km decoupled from biological sampling. The mean and SD do not include the Skyring Sound 2015 station as it shows extreme values for all variables (see Table S1). However, the values from that sample are shown in parentheses for comparison. NA: no available data.

Table 3. Relative percentages of E. huxleyi A morphotypes recorded throughout southern Patagonia fjords. Mean, standard deviation (SD), and maximum and minimum percentages of five E. huxleyi morphotypes recorded in inner surface waters of southern Patagonia (PAT; n = 883 cells counted in 23 samples) and down to 50 m in the Archipelago Madre de Dios western zone (AMD; n = 1012 cells counted in 27 samples) during the late austral spring 2015 and early spring 2017.

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3.2 Early spring southern Patagonia 2017

The hierarchical clustering based on the physical–chemical conditions in early austral spring 2017 indicated a separation between the WSM and the IC, whereas the three stations in the AMD were distributed between the two clusters (Fig. 5a). Surface salinity ranged from 25 to 30 along the south–north track, with saltier waters (>28) around the AMD (Sts. 30–32) and southward in the WSM zone (Sts. 22–24) and fresher waters in the southern IC (<27 at Sts. 25–27) (Fig. 5b). Surface temperatures ranged from a minimum of 6.3°C in the southern IC to >7.7°C in the AMD and northern IC (max. 8.1°C at St. 29, Figs. 5a–b, 54b). Surface waters were undersaturated with respect to pCO₂ at all stations (range: 320–396 µatm) except for station 26 (412 µatm). Lower Ω_{cal} levels prevailed in the southern IC (range: 1.5–1.9) with higher Ω_{cal} in the AMD zone (range: 2.1–2.4; Figs. 5b, S5b). Nitrate, silicate, and phosphate concentrations (Fig. 5c) were mostly in the range 6.0–8.3, 2.2–5.4, and 0.3–0.7 µm, respectively, with the highest levels of nitrate (9.5–12.8 µm) and phosphate (0.8–1.0 µm) at Sts. 22–23, 26, and 30, whereas stations 30 and 26 registered the highest DSi (12 and 7 µm, Fig. 5c, Table 2). Higher photosynthetic biomass (chl a > 2.7 µg L⁻¹) was recorded around the eastern AMD (St. 29, 31–32) and WSM station 23, whereas the western AMD (St. 30), southern IC (Sts. 25–26), and WSM (St. 22) yielded the lowest measured biomass (chl a < 1.3 µg L⁻¹; Fig. 5d). Variation in chl a reflected variation in diatom biomass (Fig. 5d).

The dominant coccolithophore taxon during early spring 2017 was again *E. huxleyi* (>96%). Abundances ranged
Figure 5. Physical, chemical, and nutrient conditions, chlorophyll-α levels, carbon biomass by *E. huxleyi* and diatoms, and abundances and calcification level of *E. huxleyi* recorded in surface waters of southern Patagonia during early austral spring 2017. (a) Hierarchical clustering on temperature, salinity, pH, pCO₂, and Ω calcite surface values on 11 water samples collected for plankton analysis; (b) salinity, temperature, Ω calcite, and pCO₂ levels; (c) nitrate, dissolved silicate, and phosphate levels; (d) total chlorophyll *a* and total carbon biomass by *E. huxleyi* and diatoms and semi-quantitative estimation of diatom abundances (SEM); and (e) total abundances of *E. huxleyi* and relative abundances of three *E. huxleyi* morphotypes. All samples were taken <5 m in depth. Stations 25–26 and 30 were conducted at night. Morphotype abbreviations as in Fig. 3.

from $1.69 \times 10^4$ to $9.06 \times 10^4$ cells L$^{-1}$ (Fig. 5e). The *E. huxleyi* carbon biomass averaged $0.5 \pm 0.3$ µg C L$^{-1}$ (in 11 samples), reaching both maximal and minimal values (0.2 and 1 µg C L$^{-1}$, respectively) in the southern IC (Sts. 25–27). In contrast, the opal-derived diatom carbon biomass averaged $40 \pm 17$ µg C L$^{-1}$, with lower values (<18 µg C L$^{-1}$) in the AMD at St. 30 and IC at St. 25 (Fig. 5d). While fixed samples for standard microplankton analysis were not available, large chains of *Skeletonema* spp., *Thalassiosira* spp., and *Chaetoceros* spp. were noted as frequent in samples observed by SEM (Fig. S8) and were likely significant contributors to opal. Similar to the 2015 survey, the moderate A morphotype dominated the *E. huxleyi* assemblages along the 2017 track (Fig. 5e). Cells of the lightly calcified A morphotype were sporadically observed, whereas the highly calcified A-CC and R/hyper-calcified morphotypes were again restricted to the AMD zone (Fig. 6, Table S1; note the low abundances of the R/hyper-calcified morphotype were only detected at other depths, so they do not appear in Fig. 5e).

Two CTD profiles were performed in the AMD zone: one in the western limestone AMD basin (St. 30) and another profile southwest of Escríbano Island at the eastern silicate AMD basin (St. 32) (Fig. 6a, b). The profiles covered the euphotic zone (down to 27 m in St. 32; St. 30 was conducted at night) as well as subsurface layers (25–75 m depth). In both stations, temperature and salinity increased with depth, with maximum density stratification between 5–10 m. In the western AMD profile, Ω$_{calc}$ was low at the surface (2.1) due to the lowest salinity, but rose to a maximum at 5 m due to a minimum in pCO₂ and increasing salinity, whereas below 20 m, pCO₂ rose and Ω$_{calc}$ dropped (Fig. S9e). At the eastern AMD site, in contrast, Ω$_{calc}$ increased with depth despite increasing pCO₂ (Fig. S9f). At both stations, photosynthetic biomass was mainly confined to the upper 25 m of the water column, with chl-*a* peaks at 5 and 10 m (0.7 and 3.1 µg L$^{-1}$, respectively) and dropping close to zero below 40 m in the western AMD, while remaining near 1 µg L$^{-1}$ even at depths $\geq$ 50 m in the eastern AMD (Figs. 6a–b, S9a–b). The eastern AMD zone exhibited higher chl *a* and bSi when compared to the western AMD, despite no depletion of phosphate, nitrate, and DSi being observed at either site (Figs. 6a–b, S9a–d).

At both sites, *E. huxleyi* dominated the coccolithophore assemblages across all depths, with another six coccolithophore species observed in subsurface AMD waters (Table S3). *E. huxleyi* and diatom abundances were highest in the surface at both sites (Fig. 6c–d). However, reflecting the chl-*a* profile, estimated diatom biomass remained relatively high at depth compared to surface values (dropping by only about 40%), and *E. huxleyi* abundance and biomass also dropped less with depth in the eastern AMD compared to the western AMD (Fig. 6c–d). In both stations, the composition of *E. huxleyi* morphotypes was similar at all depths, character-
Emiliania huxleyi assemblages.

lightly calcified and R/hyper-calcified morphotypes were either undetected or represented a minor fraction of coccolithophore species (i.e., G. ericsonii, G. muellerae, G. parvula), as well as the diatoms Lioloma spp., pennate diatoms (<50µm length), Nitzschia spp., Pseudo-nitzschia cuspidata, and Asteromphalus sarcophagus.

The two first axes of the CA accounted for 60% of the total explained variability and indicated that the highest E. huxleyi and low diatom biomasses were associated with increasing temperatures (Fig. 8). Intermediate E. huxleyi biomasses were associated with high diatom biomasses and increasing gradients of salinity, pH, and $\Omega_{\text{cal}}$, whereas low E. huxleyi biomasses were associated with intermediate diatom biomasses and increasing pCO$_2$. However, none of the considered environmental variables had a significant fit in the envfit test.

3.4 Niche analysis of Emiliania huxleyi morphotype responses to environmental conditions

The OMI analysis depicted differences in the realized niches of the E. huxleyi morphotypes throughout Patagonian fjords in 2015 and 2017 (Fig. 9a, Table S6). The OMI plot showed station 15 from 2015 as an outlier, characterized by extremely low salinity and high pCO$_2$. The OMI axis 1 (91.02% of explained variability) was negatively related to $\Omega_{\text{cal}}$, whereas the OMI axis 2 (8.42% of explained variability) was positively related to salinity and pH and negatively related to temperature and pCO$_2$. The envfit test indicated that all variables had a significant fit ($R^2 > 0.88$, $p < 0.01$; Table S7). Only the moderate A and R/hyper-calcified morphotypes showed significant OMI values ($p < 0.05$, Table S6). The moderate A morphotype was the most generalist (OMI = 0.07, Tol = 1.23), observed in all samples (except St. 15 in 2015). The R/hyper-calcified morphotype, observed exclusively in the AMD zone, was the most specialized morphotype (OMI = 4.77, Tol = 0.75). The A-CC morphotype (OMI = 1.43, Tol = 1.68), observed in the AMD and northern IC, showed intermediate habitat preferences (Fig. 9a), but the OMI for this morphotype did not meet the threshold for significance ($p = 0.060$). The RTol for the R/hyper-calcified morphotype was 12% (Table S6), indicating that most variability in its realized niche was accounted for by the environmental variables included in the analysis.

The extended domain OMI analysis (Fig. 9b, Table S8) indicated a clear separation between the Patagonian fjords and coastal and oceanic waters off central and northern Chile and Peru. The OMI axis 1 (74.57% of explained variability) was negatively related to temperature, salinity, and $\Omega_{\text{cal}}$, whereas the OMI axis 2 (25.32% of explained variability) was positively related to pH and negatively related to pCO$_2$. The envfit test indicated that all variables had a significant fit ($R^2 > 0.88$, $p < 0.01$; Table S9). All coccolithophore species and E. huxleyi morphotypes showed

3.3 Emiliania huxleyi abundance vs. diatoms

The nMDS depicted a clear separation between the Patagonian fjords and the oceanic/coastal areas regarding the composition of coccolithophorid and diatom assemblages (Fig. 7). The IndVal analysis (Table S5) identified only the E. huxleyi moderately calcified morphotype as an indicator of the fjord locations, along with the diatoms Thalassiosira spp., Stephanopyxis turris, Leptocylindrus spp., and Chaetoceros spp. The coastal/oceanic locations were more characterized by the lightly calcified and A-CC morphotypes and the other coccolithophore species (i.e., G. ericsonii, G. muellerae, G. parvula), as well as the diatoms Lioloma spp., pennate diatoms (<50µm length), Nitzschia spp., Pseudo-nitzschia cuspidata, and Asteromphalus sarcophagus.

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Figure 7. (a) Non-metric multidimensional scaling (nMDS) based on coccolithophore and diatom abundances attained in southern Patagonia fjords during late spring 2015 (this study) and other coastal/oceanic areas (data from von Dassow et al., 2018). (b) Heatmap showing abundances of coccolithophore and diatom species used in the nMDS. The horizontal dendrogram (based on the nMDS sample scores) shows a clear separation between Patagonia fjords (red cluster) and coastal/oceanic (blue cluster) areas, whereas the vertical dendrogram (based on the nMDS species scores) indicates the separation of species in two main clusters. Black and blue species labels depict species with significant values in the IndVal analysis. A-CC: Emiliania huxleyi A-CC morphotype; Amphi: Amphiprora spp.; Asar: Asteromphalus sarcophagus; Aster: Asteromphalus spp.; Astp: Asterionellopsis spp.; Cen.la: centric diatoms >100 µm diameter; Cen.me: centric diatoms 40–100 µm diameter; Cen.sm: centric diatoms <40 µm diameter; Chae: Chaetoceros spp.; Clep: Calcidiscus leptoporus; Coret: Corethron spp.; Cer/Gui: Cerataulina spp. + Guinardia spp.; Deton: Detonula spp.; Dityl: Ditylum spp.; Eucam: Eucampia spp.; Geri: Gephyrocapsa ericsonii; Gmue: Gephyrocapsa muellerae; Gpar: Gephyrocapsa parvula; LC: Emiliania huxleyi lightly calcified morphotype; Lepto: Leptocylindrus spp.; Licmo: Licmophora spp.; Liol: Lioloma spp.; MC: Emiliania huxleyi moderately calcified morphotype; Nit.la: Nitzschia spp. large >100 µm length; Nit.me: Nitzschia spp. medium 100–50 µm length; Nit.re: Nitzschia reversa; Nit.sm: Nitzschia spp. <50 µm length; Pcus: Pseudo-nitzschia cuspidata; Pen.la: pennate diatoms >100 µm length; Pen.me: pennate diatoms 50–100 µm length; Pen.sm: pennate diatoms <50 µm length; Pla.so: Planktoniella sol; Plagi: Plagiogrammopsis spp.; Pro.al: Proboscia alata; Pse.ni: Pseudo-nitzschia spp.; R/h: Emiliania huxleyi R/hyper-calciﬁed morphotype; Rhizo: Rhizosolenia spp.; Skele: Skeletonema spp.; Ste.tu: Stephanopyxis turris; Stria: Striatella spp.; Thala: Thalassiosira spp.; Thaln: Thalassionema spp.

4 Discussion

4.1 Patagonian coccolithophore communities dominated by E. huxleyi

Emiliania huxleyi was the only coccolithophore widely distributed along the fjords and inner channels of southern Patagonia and always represented >96% of total coccolithophore abundance and >89% of coccolithophore biomass, during both early and late spring. The low diversity of coccolithophore assemblages, dominated by E. huxleyi, is a common spring–summer feature in both the Patagonian and Norwegian fjord systems. In the case of southern Patagonia, the neighbouring Pacific has higher diversity (Beaufort et al., 2008; Menschel et al., 2016; von Dassow et al., 2018), but the Southern Ocean assemblages also show low diversity dominated by E. huxleyi (Cubillos et al., 2007; Saavedra-Pellitero et al., 2014; Charalampopoulou et al., 2016; Saavedra-Pellitero et al., 2019). The low diversity
in southern Patagonian waters thus may partly reflect this latitudinal trend, although more detailed seasonal studies, including sampling along vertical profiles, might reveal significant additional coccolithophore diversity in the Patagonian and Norwegian fjords.

4.2 Abundance of *E. huxleyi* in Patagonia compared to nearby oceans

During early and late spring, standing stocks of *E. huxleyi* in the Patagonian fjords and inner seas were moderate compared with those documented in nearby coastal and oceanic regions and within the range of background stocks reported in the Norwegian fjords and North Sea (Table 4 and references therein). The high *E. huxleyi* abundances typical of spring blooms in the Norwegian fjords were not observed in either early or late spring in the present study despite the similar temperature, salinity, and Ω̅cal conditions in both fjord systems. No *E. huxleyi* blooms have been reported in Patagonian fjords, although this might be due to limited observations and methodological issues. For example, many phytoplankton studies in the area (e.g., Alves-de-Souza et al., 2008; González et al., 2013) as well as standard phytoplankton monitoring in the zone (Vivanco and Seguel, 2009) often rely on samples fixed with Lugol’s solution acid, which would not preserve coccoliths, or the studies have only focused on larger phytoplankton size classes (e.g., Paredes et al., 2014). Shallow water depth and frequent cloud cover limit satellite observations of particulate inorganic carbon (PIC) within the

Figure 8. Correspondence analysis (CA) assessing the relationship between *E. huxleyi* and diatom biomasses converted to categoric values (i.e., low, intermediate, and high biomasses) in Patagonia fjords during late spring 2015 and early spring 2017. The envit function of the vegan package (R software) was used to fit the environmental variables to the CA plot (no variable was significant; p > 0.05).

4.3 Variation in *E. huxleyi* with environmental factors

It has been previously proposed that the realized niche of *E. huxleyi* is partly defined by physical and chemical conditions unfavourable to large diatoms (Tyrrell and Merico, 2004; Smith et al., 2017). During late spring the Patagonia fjords, *E. huxleyi* reached higher abundances in the southern IC when the temperature was above 8 °C and macronutrients and larger diatoms were the lowest, consistent with the pattern previously reported more generally for nanophytoplankton based on size-fractionated chl *a* for this geographic area (e.g., Cuevas et al., 2019). However, the CA showed that the lowest levels of *E. huxleyi* were associated with intermediate levels of larger diatoms, and intermediate levels of *E. huxleyi* were associated with the highest levels of larger diatoms, suggesting a unimodal relationship between these two planktonic groups, possibly affected by factors not assessed in this study, such as nutrient supply and mixed-layer depth (Margalef, 1978; Cermeño et al., 2011), or predation (Nejstgaard et al., 1997).

The Ω̅cal—the saturation state of calcite, a parameter often assumed to constrain calcification (e.g., Zondervan et al., 2001; Kleypas et al., 2006; but see Cyronak et al., 2016) – was subject to large spatial variations in surface waters, from relatively high Ω̅cal levels in the AMD zone (range: 2.1–3.6), to moderate Ω̅cal in the interior WSM, low Ω̅cal in the southern IC (range: 1.5–2.2), and subsaturation in the SS (0.5). The range of surface Ω̅cal recorded along southern Patagonia was comparable to that reported for the Norwegian seas (Jones et al., 2019). Whereas the highest Ω̅cal values observed at the AMD were not as high as those observed in the global ocean (Takahashi et al., 2014), the lower values at the southern IC were comparable to values reported previously (range: 1.8–2.8) from high-CO₂-upwelling conditions in central and northern Chile (Beaufort et al., 2008, 2011; von Dassow et al., 2018). While low surface Ω̅cal at coastal waters of northern and central Chile are related to the upwelling of high-pCO₂-DIC subsurface waters, the freshening (and associated drop in DIC, salinity, and Ca²⁺ caused by dilution) and latitudinal—seasonal cooling (enhancing CO₂ solubility) have major roles in lowering CO₂− and Ω̅cal in southern Patagonia. These contrasting systems offered the possibility to observe whether the ecological trends related to low Ω̅cal depend on context.
4.4 Comparison of *E. huxleyi* morphotypes in Patagonia to nearby oceans and Norwegian fjords

There was some variability in the vertical distribution of the *E. huxleyi* morphotypes in the water column. The lightly calcified coccoliths appeared associated with subsurface waters in both seasons sampled at the locations, so they might be associated with intrusion of these waters. However, the samples within the euphotic zone were generally similar to each other within a given sample station. Thus, for the purposes of the questions in this study, the use of surface samples to describe morphotype distributions is expected to be reasonable, but use of vertical profiles might have permitted a higher ability to explain variability in the OMI analysis.

The *E. huxleyi* populations in the Patagonian fjords were morphologically distinct from surrounding coastal or open ocean populations in the eastern South Pacific, the Southern Ocean, and the Atlantic. The Atlantic Patagonian Shelf *E. huxleyi* populations are reported to be dominated by B/C morphotypes (Poulton et al., 2011, 2013). Southern open ocean populations of *E. huxleyi* are dominated by B morphotypes (including the B, B/C, C, and O types; Saavedra-Pellitero et al., 2014; Saavedra-Pellitero et al., 2019), and A morphotypes were reported to represent only a small fraction. However, C and O morphotypes were very rare in Patagonian inland waters, and B and B/C morphotypes were undetected. Although the moderately calcified and robustly calcified A morphotypes have also been shown to be present in eastern South Pacific coastal and open ocean waters (von Dassow et al., 2018), the dominance of these A morphotypes was particular to Patagonian interior waters. This conclusion was supported by the IndVal analysis (Table S5), where moderately calcified and robustly calcified A morphotypes were consolidated for final statistical analyses as they are not easily distinguished by objective morphological characters and were present in all samples, and preliminary analysis revealed completely overlapping realized niches. Both the moderately calcified and robustly calcified A morphotypes are also observed as dominant in the Norwegian fjords (Table 4) (Young, 1994). The lightly calcified A morphotype was rare and did not show any clear pattern in its distribution. The A-CC morphotype has been associated with coastal upwelling zones in the Atlantic (Giraudeau et al., 1993; Smith et al., 2012; Henderiks et al., 2012) but not reported from the
Norwegian fjords or the Southern Ocean. In both early and late spring, R/hyper-calcified and A-CC *E. huxleyi* appeared only at the Pacific border of southern Patagonia (AMD zone). Thus, *E. huxleyi* populations of both Patagonian fjords and Norwegian fjords share a similar morphotype composition.

4.5 Niche analysis of *E. huxleyi* morphotypes related to carbonate chemistry conditions

The broader niche breadth by the moderately calcified A morphotype contrasted with the marginal niche of the R/hyper-calcified forms in Patagonia (Fig. 9a). The lightly calcified A morphotype also showed a low tolerance (more specialist), but this was not statistically significant. In order to extend the realized niches derived in Patagonia, we complemented the OMI analysis with a sample set of nearby oceanic and coastal sites (data from von Dassow et al., 2018), in some of which the moderately calcified A morphotype, unlike in Patagonia, was less abundant than other *E. huxleyi* morphotypes and coccolithophore species. According to OMI analysis, the niche differentiation along Patagonia is mostly driven by the pH–Ω_{calc} conditions, but temperature and salinity conditions also become important. In this extended domain, both the moderately calcified A morphotype and the A-CC morphotype appeared to be generalists, with high Tol values (Fig. 9b). The lightly calcified morphotype also appeared to be a generalist in the extended domain. However, we caution that while the lightly calcified *E. huxleyi* was almost exclusively the lightly calcified A morphotype in Patagonia, there was a continuum of lightly calcified A, B, and C morphotypes (and some lightly calcified cells were difficult to classify among these types) in some of the coastal and oceanic sites. Proper differentiation between B, C, and C based on coccolith length would require strict morphometrics, which we did not perform due to the difficulty in accurate measurements on full coccospheres.

### Table 4. Comparison of *E. huxleyi* standing stocks and morphotypes recorded in northern and Patagonian fjord systems and nearby coastal/ocean locations. Temperature and salinity ranges are shown. Only data < 10 m in depth were included. ESP: eastern South Pacific; SO: Southern Ocean; APS: Atlantic Patagonian Shelf; SU: summer; SP: spring; WI: winter; AU: autumn; NA: no available data.

| Location | Concentration cells × 10^{3} L^{-1} | *E. huxleyi* coccoliths | Main morph | Year/season | Temp. range | Salinity range | References |
|----------|-------------------------|-----------------------|------------|-------------|--------------|---------------|------------|
| **Northern Hemisphere fjords** | | | | | | | |
| Oslo fjord | 24–36 000^{b} | NA | NA | 1939/SU | 8.2–19.6 | 8.2–28.4 | Birkenes and Braarud (1952) |
| Western coast and fjords | 250–115 000^{b} | NA | NA | 1955/SU | 11–15 | 28–33 | Berge (1962) |
| Oslo fjord | 385–14 500^{b} | NA | NA | 1981/SU | 16–20 | 20–21 | Paasche and Kristiansen (1982) |
| Bokna fjord | 3000–4000^{b} | NA | NA | 1981/SU | 11–13 | 28–30 | Era (1989) |
| Sammanger fjord | < 10–7000^{a} | NA | A | 1992/SU | 8–10 | 10–30 | Kristiansen et al. (1994); Young (1994) |
| Fauskangerpollen, Nordåsvannet | 1–28 000^{a} | NA | NA | 1993/SP | 7–16 | 15–30 | Fernández et al. (1996) |
| **Nearby coastal and oceanic zones** | | | | | | | |
| Norwegian Sea | 200–3000^{d} | dominate | NA | 1987–1992 | 6–10 | > 35 | Samtleben et al. (1995) |
| Norwegian Sea | < 10–3000^{d} | dominate | NA | 1987–1995 | 6–10 | < 35 | Baumann et al. (2000) |
| Outer Oslo fjord, Skagerakk | all year round^{e} | NA | NA | 2009–2011 | 2–16 | 22–33 | Egge et al. (2015) |
| Outer Oslo fjord, Skagerrak | 171–254^{f} | NA | A | 2013/SU | 18.5 | 23.7 | Gran-Stadniczeñko et al. (2017) |
| Northeast Atlantic and North Sea | 100–1500^{f} | dominate | A, B | 1990/SU | NA | NA | van Heijningen et al. (1991) |
| North and Norwegian seas | < 3–900^{f} | 40–100 | A^{i} | 2008/SU | 11–18 | 30–35 | Charalampopoulos et al. (2011) |
| **Patagonia fjords** | | | | | | | |
| W. Magellan Strait | ∼ 52–54^{h} | S | ~ 100^{h} | dominate | NA | 1991/SU | NA | NA | Zingone et al. (2011) |
| Fjords and channels | ∼ 50–54^{h} S | 12–276^{h} | > 98 | A | 2015/SP | 7.1–10.0 | 26.6–31.3 | This study |
| Fjords and channels | ∼ 50–54^{h} S | 17–91^{h} | > 96 | A | 2017/SP | 6.3–8.1 | 25.1–29.8 | This study |
| **Nearby coastal and oceanic zones** | | | | | | | |
| ESP along | ∼ 16–20^{d} | S, 73–80^{d} W | 1–34^{d} | 8–99 | A | 1964/SU | 21.9–24.9 | NA | Hagino and Okada (2006) |
| ESP along | ∼ 30–33^{d} | S | 60 – 90 | A, R^{f} | 2004/SP | 14.8–19.8 | 34.1–35.1 | Beaufort et al. (2008, 2011) |
| ESP along | ∼ 35–38^{d} | S, 73–78^{d} W | 12–134^{d} | > 50 | A, R^{f} | 2004/SP | 10.7–15.3 | 33.4–34.6 | Menschel et al. (2016) |
| ESP along | ∼ 12^{d} | S, 77–78^{d} W | 1–531^{d} | > 76 | A, R^{f} | 2014–15/AU,SP | 14–21 | 35–35.2 | Alvies et al. (2016) |
| ESP along | ∼ 13–33^{d} | S, 70–86^{d} W | 1–76^{d} | 30–100 | A, R^{f} | 2011,13/SP, WI | 12.3–19.8 | 33.8–35.5 | von Dassow et al. (2018) |
| SO Pacific | ∼ 45–55^{d} S, 80–100^{d} W | NA | 50–100 | C | 1967–1970 | 1–14 | NA | Melhynes et al. (1970) |
| SO Pacific | ∼ 54–58^{d} S, 80–97^{d} W | 31–61^{d} | dominate | C, B/C | 2009/SP | 4.3–5.7 | ∼ 3.4 | Saavedra-Pellitero et al. (2014) |
| SO Drake Passage | ∼ 57–62^{d} S | 1–580^{d} | > 80 | B/C | 2009/SP | 2–10 | 33.7–34.0 | Charalampopoulos et al. (2016) |
| SO Drake Passage | ∼ 57–62^{d} S | 1–214^{d} | dominate | B/C | 2016/SU | 2–10 | 33.3–34.0 | Saavedra-Pellitero et al. (2019) |
| APS along | ∼ 38–54^{d} S, 51–63^{d} W | 180–3006^{d} | dominate | B/C | 2008/SP | 7–18 | 33.4–35.6 | Poulton et al. (2011, 2013) |

* *E. huxleyi* as a component of larger coccolithophore assemblages. When information about other coccolithophores was provided, the percentage abundance of *E. huxleyi* was dominant. "NA" means no information was provided for the parameter. \(^{b}\) Estimated through sedimentation chamber and inverted microscopy. \(^{c}\) Estimated by Palme Maloney chamber and inverted microscopy. \(^{d}\) Estimated through filtration and scanning electron microscopy analysis. \(^{e}\) Estimated through high-throughput sequencing and operational taxonomic unit (OTU) analysis. They find *E. huxleyi* throughout the year but concentrated between summer–autumn. \(^{f}\) Cultured strains were identified at morphotype level by using immunofluorescence assay, and abundances were estimated through scanning electron microscopy analysis. \(^{g}\) Estimated through filtration and cross-polarized microscope. \(^{h}\) Estimated using an automated coccosphere recognition software. \(^{i}\) The “characteristic” *E. huxleyi* morphotype showed in Fig. 4. \(^{j}\) The R/hyper-calcified morphotype dominating the *E. huxleyi* assemblages in neritic zones.
of less common morphotypes, especially in low-abundance populations (as coccospheres may lack coccoliths in a correct orientation for accurate measurement). Thus the generalist behaviour of lightly calcified morphotypes in the OMI analysis that combined fjord, coastal, and open ocean sites is likely an artefact. We suspect that lightly calcified A, B, B/C, and C morphotypes might actually each exhibit specialist behaviours in distinct but overlapping niches. In fact, a laboratory study reported that B/C morphotype strains only calcified substantially in a relatively narrow range of carbonate conditions (Müller et al., 2015). In contrast, the very distinct R/hyper-calcified morphotype exhibited restricted preferences in terms of $\Omega_{\text{cal}}$, temperature, and salinity but a broad niche in terms of $\text{CO}_2$ and pH (Fig. 9b).

The R/hyper-calcified morphotype, in which there is both fusion of distal shield elements and closure or partial closure by overcalcification of the central area, has so far only been reported as prevalent in the high-$\text{CO}_2$ and low-pH upwelling zone of the eastern South Pacific (Beaufort et al., 2011; Alvites, 2016; von Dassow et al., 2018), although it has been seen (and reported as rare) in both Australian waters (Cubillos et al., 2007) and the Drake Passage (Saavedra-Pellitero et al., 2019). Experimental findings that the R/hyper-calcified morphotype did not perform better than the moderately calcified A morphotype under high $\text{CO}_2$/low pH/low $\Omega_{\text{cal}}$ (von Dassow et al., 2018) might be explained by the OMI analysis suggesting a possible narrow unimodal response to $\Omega_{\text{cal}}$, which would not have been detected in the experiments of von Dassow et al. (2018), where $\Omega_{\text{cal}}$ values of 1.4 vs. 3.3 were tested in the lab. The studies of Langer et al. (2009) and Müller et al. (2015) did find that R morphotype strains did seem more resistant to high $\text{CO}_2$ and low pH than other A morphotypes, either in growth rate or in PIC production. Those studies used either 4-fold higher light levels (Langer et al., 2009) or continuous light (Müller et al., 2015), and low light has been shown to increase the sensitivity to OA, specifically of an R morphotype strain (Rokitta and Rost, 2012), highlighting that the R/hyper-calcified morphotype might be selected by interactions with other variables. Alternatively, the R/hyper-calcified morphotype might be selected by an unidentified condition particular to the southeastern Pacific that correlates with the $\Omega_{\text{cal}}$, temperature, and salinity of its realized niche.

The present study shows that the OMI analysis can be useful for identifying how parameters may determine the realized niches of both species and genetically determined phenotypic variants within a species. For example, in the extended domain (Fig. 9b), $\Omega_{\text{cal}}$, temperature, and salinity were important in defining the narrowness vs. breadth of niches among the $E$. huxleyi A morphotype, $E$. huxleyi R/hypercalcified morphotype, and closely related $G$. parvula species, while the habitat centres (niche positions) of the different $E$. huxleyi morphotypes lined up approximately on a gradient of pH vs. $\text{CO}_2$. Calcification rate has been demonstrated to vary among $E$. huxleyi strains according to maximum photosynthetic rate but also $\text{CO}_3^{2-}$ concentration at their sites of origin, while maximum photosynthetic rate also varied among the strains with $\text{CO}_2$ at the site of origin (Rickaby et al., 2016). Several recent studies have called into question the importance of $\Omega_{\text{cal}}$ in determining the response of calcifying organisms, with both theoretical and laboratory measurements support that the concentrations of $\text{HCO}_3^-$, $\text{CO}_3^{2-}$, and $\text{H}^+$ are more important (Kottmeier et al., 2016; Bach et al., 2015; Gafar et al., 2018). However, while $\text{HCO}_3^-$ and $\text{CO}_3^{2-}$ were formally excluded from the OMI analyses as redundant variables, these vary more with $\text{CO}_2$ and pH, respectively, which formed an axis along which the niche of the R/hyper-calcified morphotype was broad, rather than the orthogonal gradient in $\Omega_{\text{cal}}$. We caution that calcification rate might not relate in a simple way to the morphotypes observed here. Nevertheless, such trade-offs offer crucial clues into how traits related to calcification may be selected by the environment, resulting in the environmental patterns observed here. It might be worth exploring experimentally whether there is a role for $\Omega_{\text{cal}}$ separate from other carbonate parameters in selecting the R/hypercalcified morphotype.

The OMI analysis presented here was limited as we focused mostly on the carbonate system, and this may be reflected in that sometimes half of the total variability was not explained by included variables. As mentioned above, several studies have shown that light can impact sensitivity of $E$. huxleyi to OA (Rokitta and Rost, 2012; Jin et al., 2017), although the effect reported differs in some studies (Zhang et al., 2019). The trade-offs in energy and C balances reported by Rickaby et al. (2016) would suggest that light and mixed-layer depth might also be important parameters to consider in future studies. Nutrient limitation can also modify calcification, although a careful chemostat study suggested that nutrient and $\text{CO}_2$ impacts were independent (Müller et al., 2017). The impact of biotic factors was also not assessed with the OMI analysis here, though it is still not clear whether coccoliths might play roles in defence against either grazing or viruses (Harvey et al., 2015; Strom et al., 2018; Johns et al., 2019; Haunost et al., 2020).

A striking result from the OMI analysis was that all the $E$. huxleyi morphotypes, even the more specialized R/hypercalcified type, exhibited much greater niche breadth (higher Tol values) than the other coccolithophore species. The three $G$. parvula and $G$. ericsonii showed Tol values that were more than 10-fold lower than the most specialist $E$. huxleyi morphotypes. The small $G$. parvula and $G$. ericsonii showed Tol values that were more than 10-fold lower than the most specialist $E$. huxleyi morphotype. Despite the evidence for a genetic underpinning of $E$. huxleyi morphotypes (Krueger-Hadfield et al., 2014), as well as evidence of a high level of genomic content variability in $E$. huxleyi (von Dassow et al., 2015), phylogenetic and phylogenomic evidence does not clearly support for it
to be split into different species (Bendif et al., 2016; Filatov, 2019). If the ubiquitous taxon is less susceptible to environmental change compared to marginal taxa (i.e., marginality or richness vs. tolerance are inversely correlated; Dolédé et al., 2000; Hernández et al., 2015), the exceptional generalist behaviour exhibited by *E. huxleyi* compared to other coccolithophores suggests it may be more plastic and more adaptable in the face of environmental change.

The lower values of pH and $\Omega_{\text{cal}}$ observed here approached levels predicted for higher latitudes of the global ocean at the end of the century under high-emission scenarios such as RCP8.5 (Feely et al., 2009; Hartin et al., 2016). Our results suggest that ongoing changes in ocean chemistry may result in decreases in coccolithophore diversity, leading to more numerical dominance of *E. huxleyi* compared to other coccolithophores, as well as decreased phenotypic diversity within *E. huxleyi*. The extended-domain niche analysis in the present study would suggest that *C. leptoporus* might be less adaptable than *E. huxleyi*. Some studies found that *C. leptoporus* was relatively resistant to OA in the lab (Langer et al., 2006), but others reported that it is sensitive and that its PIC / POC (particulate organic carbon) ratio, considered important in determining ballast effects, is especially negatively affected by OA. Species such as *C. leptoporus* can be much more important than *E. huxleyi* in carbonate export due to their production of much heavier coccolithophores which sink faster and dissolve more slowly (e.g., Ziveri et al., 2007; Menschel et al., 2016; Menschel and González, 2019). Thus, a change to more *E. huxleyi*-dominated coccolithophore communities might negatively impact carbonate export.

5 Conclusions

Our study of how *E. huxleyi* abundances and morphotypes respond to the highly dynamic physical and chemical environments of southern Patagonia yielded seven principal findings.

1. The only coccolithophore that was a regular and ubiquitous component of the phytoplanktonic assemblages throughout the surface waters of the southern Patagonian fjords/channels was *E. huxleyi*. It occurred under a wide range of carbonate chemistry conditions and was only absent in the Skyring Sound zone where $\Omega_{\text{cal}} < 1$.

2. Although *E. huxleyi* never reached more than a small fraction of total plankton carbon biomass (< 13% of nano- and microplankton assemblages counted by microscopy), it reached moderate abundances (range: 12–276 × 10³ cells L⁻¹) comparable to adjacent coastal and oceanic areas, and within the lower range of stocks reported from Norwegian fjords (1–115 000 × 10³ cells L⁻¹).

3. *Emiliania huxleyi* abundance was highest (> 100 × 10³ cells L⁻¹) when assemblages of large diatoms were lowest (< 10 × 10³ cells L⁻¹), in late spring waters with lower macronutrients, consistent with it being most important in the absence of large diatoms.

4. In terms of morphotypes, the *E. huxleyi* populations in the southern Patagonian fjords/channels were similar to Norwegian fjords (dominated by moderately calcified and robustly calcified A morphotype) and very distinct from populations previously documented in the Southern Ocean and Drake Passage and the Patagonian Shelf of the Atlantic (where C or B/C morphotypes were reported as dominant) and from the eastern South Pacific coastal upwelling zone, where the R/hyper-calcified morphotype dominated.

5. Niche analysis shows that the moderate A morphotype and A-CC morphotypes are generalists, whereas the R/hyper-calcified morphotype has a more marginal (specialized) realized niche.

6. The association of the R/hyper-calcified morphotype to high $\Omega_{\text{cal}}$ in southern Patagonia, where $\Omega_{\text{cal}}$ is driven principally by freshwater input, contrasts with its dominance in the upwelling system of central Chile to Peru, where low $\Omega_{\text{cal}}$ is due to high CO₂. This morphotype occupies a narrow range of $\Omega_{\text{cal}}$ values compared to the A-CC and moderate A morphotypes.

7. The moderate A, A-CC, and R/hyper-calcified *E. huxleyi* morphotypes all display higher niche breadth (more generalist behaviour) than closely related coccolithophores, suggesting that *E. huxleyi* may be ecologically more plastic and have more capacity for adaptation in the face of environmental change than other coccolithophores.

Data availability. All data resulting from this study are available from the corresponding authors upon request. The scanning electron micrograph image datasets can be found at https://doi.org/10.5281/zenodo.4292020 (Díaz-Rosas et al., 2021a). The inverted microscope images used for biovolume and carbon biomass estimations can be found at https://doi.org/10.5281/zenodo.5139161 (Díaz-Rosas et al., 2021b). Abundances of all plankton quantified in 2015 can be found at https://doi.org/10.1594/PANGAEA.936506 (Díaz-Rosas et al. 2021c). Coccolithophore relative abundances can be found at https://doi.org/10.1594/PANGAEA.936509 (Díaz-Rosas et al. 2021d). Physical–chemical parameters associated with abundances and biovolumes of distinct *E. huxleyi* morphotypes can be found at https://doi.org/10.1594/PANGAEA.936505 (Díaz-Rosas et al. 2021e).
Sample availability. Material for SEM characterization (filter sections) are in Dr. Peter von Dassow’s laboratory. laboratory and can be requested.

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Author contributions. FDR (conceptualization, investigation, data curation, formal analysis, writing – original draft preparation, writing – review and editing) led the study, carried out sampling in the 2015 survey, carried out light and SEM microscopic analysis, conducted characterization of microplankton and E. huxleyi assemblages and morphotype composition, performed bioluminal analysis, analyzed the relationships between E. huxleyi and environmental and biological variables, and wrote the first drafts of the paper. PvD (conceptualization, funding acquisition, supervision, validation, visualization, writing – original draft preparation, writing – review and editing) trained and supervised FDR in E. huxleyi assemblage quantification and characterization, guided analysis strategies, performed extensive re-writes of the text and figures, and provided continuous insights into interpretation of results and how to structure the manuscript. CAaD (formal analysis, supervision, validation, visualization, writing – review and editing) trained and supervised FDR in microplankton light-microscopy qualitative and quantitative analysis; performed nMDS, CA, and OMI niche analysis; guided interpretations of these results; and helped with extensive re-writes of the text and figures. EA (investigation, formal analysis, writing – review and editing) performed the analysis of carbonate system parameters, nutrients, opal and chl a and helped characterize the physical environments. EM (investigation, formal analysis, writing – review and editing) performed the analysis of carbonate system parameters, nutrients, opal and chl a and helped characterize the physical environments. EM (investigation, formal analysis, writing – review and editing) carried out sampling and CTD deployment during the 2017 survey and together with HG (funding acquisition, writing – review and editing) provided insights into the interpretation of oceanographic results. All co-authors provided key comments and editing of the final draft of the paper.

Competing interests. The authors declare that they have no conflict of interest.

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