



# Biogenic silica production and diatom dynamics in the Svalbard region during spring

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**Abstract.** Diatoms are generally the dominant contributors to the Arctic Ocean spring bloom, which is a key event in regional food webs in terms of capacity for secondary production and organic matter export. Dissolved silicic acid is an obligate nutrient for diatoms and has been declining in the European Arctic since the early 1990s. The lack of regional silicon cycling information precludes understanding the consequences of such changes for diatom productivity during the Arctic spring bloom. This study communicates the results from a cruise in the European Arctic around Svalbard, which reports the first concurrent data on biogenic silica production and export, export of diatom cells, the degree of kinetic limitation by ambient silicic acid, and diatom contribution to primary production. Regional biogenic silica production rates were significantly lower than those achievable in the Southern Ocean and silicic acid concentration limited the biogenic silica production rate in 95 % of samples. Compared to diatoms in the Atlantic subtropical gyre, regional diatoms are less adapted for silicic acid uptake at low concentration, and at some stations during the present study, silicon kinetic limitation may have been intense enough to limit diatom growth. Thus, silicic acid can play a critical role in diatom spring bloom dynamics. The diatom contribution to primary production was variable, ranging from < 10 % to ~ 100 % depending on the bloom stage and phytoplankton composition. While there was agreement with previous studies regarding the export rate of diatom cells, we observed significantly el-

evated biogenic silica export. Such a discrepancy can be resolved if a higher fraction of the diatom material exported during our study was modified by zooplankton grazers. This study provides the most direct evidence to date suggesting the important coupling of the silicon and carbon cycles during the spring bloom in the European Arctic.

## 1 Introduction

Diatoms and the haptophyte *Phaeocystis* are the dominant contributors to the Arctic Ocean spring bloom, a cornerstone event supplying much of the annual net community production (Rat'kova and Wassmann, 2002; Vaquer-Sunyer et al., 2013; Wassmann et al., 1999) that fuels Arctic food webs (Degerlund and Eilertsen, 2010, and references therein). Hydrographic and chemical changes in the Arctic water column are expected in the future, but whether these will alter diatoms' contribution to spring primary production and organic matter export remains uncertain. Some studies predict reduction in ice cover will enhance the spring bloom due to increased light availability (Arrigo et al., 2008), while others predict lower productivity driven by increased stratification and reduced nutrient supply (Schourup-Kristensen et al., 2018; Tremblay and Gagnon, 2009). Additionally, models predict that warming will lead to a shift from a diatom-dominated bloom to one increasingly dominated by flagel-

lates and picoautotrophs, which has been observed in certain sectors of the Arctic (Li et al., 2009; Lasternas and Agustí, 2010). Because the spring diatom bloom is arguably the single most important productivity event for the Arctic Ocean ecosystem (Degerlund and Eilertsen, 2010; Holding et al., 2015; Vaquer-Sunyer et al., 2013), understanding how diatoms' ecological and biogeochemical importance changes in response to system-wide physical or chemical shifts is important to predict future food web alterations. Diatoms have an obligate requirement for silicon; therefore understanding regional silicon cycling can provide insights into diatoms' activity. However, there is a current knowledge gap in regional silicon cycling, which precludes robust assessments of the spring bloom in future scenarios, e.g., Tréguer et al. (2018).

Diatom production is dependent on the availability of dissolved silicic acid ( $\text{Si}(\text{OH})_4$ ), which they use to build their shells of biogenic silica ( $\text{bSiO}_2$ ).  $[\text{Si}(\text{OH})_4]$  has been observed to be low ( $< 5 \mu\text{M}$ ) in the Norwegian Seas and declining over time (Rey, 2012). A more recent analysis demonstrated a decline in pre-bloom  $[\text{Si}(\text{OH})_4]$  concentrations by 1–2  $\mu\text{M}$  across the North Atlantic subpolar and polar regions over the last 25 years (Hátún et al., 2017). This is in stark contrast to the 10–60  $\mu\text{M}$   $[\text{Si}(\text{OH})_4]$  observed in the surface waters of the Southern Ocean and the marginal ice zone around Antarctica (Nelson and Gordon, 1982; Brzezinski et al., 2001), where  $[\text{Si}(\text{OH})_4]$  is unlikely to limit diatom growth unless iron is replete and stimulates exceptional blooms which consume Si, or assemblages are highly inefficient for Si uptake (Brzezinski et al., 2001). Additionally, the stoichiometry of  $\text{Si}(\text{OH})_4$  availability relative to nitrate ( $\text{Si} : \text{N} < 1$ ) in the source waters, which fuel the spring bloom in most of the North Atlantic and European polar seas, suggests that during a bloom cycle diatoms may experience Si limitation prior to N limitation, especially if diatoms consumed Si and N in near-equal quantities as in other diatom bloom regions (Brzezinski et al., 1997; Brzezinski, 1985; Dugdale et al., 1995).

Compared to the Southern Ocean, there is a paucity of field Si-cycling studies in the European Arctic. Reports of diatom silica production are only available from the subarctic northeast Atlantic near  $\sim 60^\circ \text{N}$ , e.g., between Iceland and Scotland (Allen et al., 2005; Brown et al., 2003), in Oslofjorden (Kristiansen et al., 2000), and in Baffin Bay (limited data; Hoppe et al., 2018; Tremblay et al., 2002); these previous studies are in zones with higher  $\text{Si}(\text{OH})_4$  availability than in the European Arctic. Other studies have reported standing stocks of  $\text{bSiO}_2$  and export in Oslofjorden or the European Arctic, e.g., Svalbard vicinity, Laptev Sea (Hodal et al., 2012; Heiskanen and Keck, 1996; Paasche and Ostergren, 1980; Lalande et al., 2016, 2013), but none have concurrent measurements of  $\text{bSiO}_2$  production. Indeed, in the last major review of the global marine silicon cycle, Tréguer and De La Rocha (2013) reported no studies with published  $\text{bSiO}_2$  production data from the Arctic.

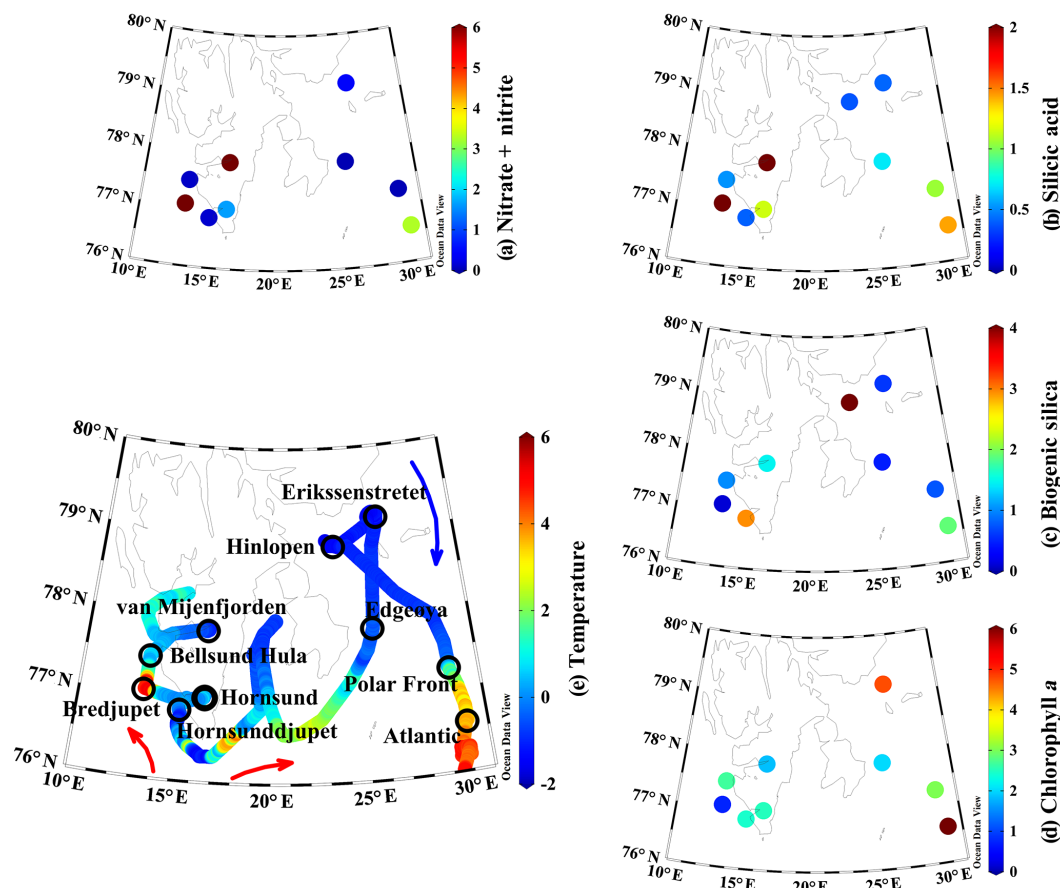
Currently, we lack a baseline understanding on diatom Si-cycling in the European Arctic and broader high-latitude North Atlantic region. And while models in the Barents Sea use Si as a possible limiting nutrient (Wassmann et al., 2006; Slagstad and Støle-Hansen, 1991), there are no field data to ground truth the modeled parameters governing diatom Si uptake. Thus, there is no contextual understanding to determine the consequences of the observed changes in regional  $[\text{Si}(\text{OH})_4]$  since the 1990s and whether these affect spring bloom dynamics. This study communicates the results from a cruise in the European Arctic around Svalbard reporting the first concurrent data sets on regional  $\text{bSiO}_2$  production and export, the export of diatom cells, and the degree of kinetic limitation by ambient  $[\text{Si}(\text{OH})_4]$ . Additionally, coupling  $\text{bSiO}_2$  production rates with contemporaneous primary production measurements provides an independent assessment for the diatom contribution to system primary production.

## 2 Methods

### 2.1 Region and sampling

This study was conducted onboard the RV *Helmer Hanssen* between 17 and 29 May 2016 as part of the broader project, ARCEX – The Research Centre for Arctic Petroleum Exploration (<http://www.arcex.no/>, last access: 29 October 2018). The main goal of this cruise was to study the pelagic and benthic ecosystem during the Arctic spring bloom around Svalbard and in the northern Barents Sea at stations which are influenced by various water masses. The cruise started in the southwestern fjords influenced by relatively warm Atlantic water, then transited east of Svalbard toward more Arctic-influenced water (Fig. 1 blue arrow) before turning south towards stations near the Polar Front and south of the Polar Front in Atlantic-influenced water (Fig. 1 red arrows).

Vertical profiles with a CTD were conducted at all stations. Hydrocasts were conducted using a Seabird Electronics 911 plus CTD with an oxygen sensor, fluorometer, turbidity meter, and PAR sensor (Biospherical/LI-CORR, SN 1060). The CTD was surrounded by a rosette with 12 5 L Niskin bottles. At two stations, Edgeøya and Hinlopen, only surface samples were collected (no vertical profiles with ancillary measurements, Fig. 1). Water was sampled from the rosette at depths within the upper 40 m (i.e., the extent of the photic layer); for any incubation described below, the approximate irradiance at the sample depth during collection was mimicked by placing incubation bottles into a bag made of neutral-density screen. Incubation bags were placed in an onboard acrylic incubator cooled with continuously flowing surface seawater. At Hinlopen, a block of ice was collected by hand within  $\sim 10$  m of the vessel and allowed to thaw in a shaded container for 24 h at ambient air temperature. After thawing, the



**Figure 1.** Surface properties during 2016 ARCEX cruise including (a) nitrate + nitrite ( $\mu\text{M}$ ), (b) dissolved silicic acid ( $\mu\text{M}$ ), (c) biogenic silica ( $\mu\text{mol Si L}^{-1}$ ), (d) chlorophyll *a* ( $\mu\text{g L}^{-1}$ ), and (e) temperature ( $^{\circ}\text{C}$ ) overlaid on station map. Station names are denoted on the map and colored arrows generalize the flow of Atlantic-influenced (red) and Arctic-influenced (blue) waters.

melted solution was homogenized and treated like a water sample for measurement of biomass and rates.

Four sediment trap arrays were deployed and collected particulate material for 19 and 23 h, depending on location. Arrays in van Mijenfjorden and Hornsund were anchored to the bottom (60 and 130 m, respectively), whereas the other two arrays (Erik Eriksenstretet, 260 m bottom depth; Polar Front, 290 m bottom depth) were quasi-Lagrangian and drifted between 14 and 16 km during the deployment. During the Erik Eriksenstretet deployment, the array was anchored to an ice floe. Arrays included sediment trap cylinders (72 mm internal diameter  $\times$  450 mm length,  $\sim$  1.8 L volume; KC Denmark) at three (van Mijenfjorden) to seven (Atlantic Station) depths between 20 and 150–200 m, based on bathymetry. After recovery, trap contents were pooled and subsampled for  $\text{bSiO}_2$  and phytoplankton taxonomy.

## 2.2 Standing stock measurements

A suite of macronutrients were analyzed at all stations except Hinlopen (just  $\text{Si(OH)}_4$ ). Water was sampled directly

from the rosette, filtered ( $0.7 \mu\text{m}$  pore size), and immediately frozen. In the laboratory, nutrients were analyzed using a Flow Solution IV analyzer (OI Analytical, USA) and calibrated with reference seawater (Ocean Scientific International Ltd., UK). Detection limits for  $[\text{NO}_3 + \text{NO}_2]$  and  $[\text{Si(OH)}_4]$  were 0.02 and 0.07 ( $\mu\text{M}$ ), respectively. No ammonium was measured. To avoid artefacts with prolonged freezing (Clementson and Wayte, 1992; Macdonald et al., 1986), samples were analyzed within 4 months of collection and standard practices were used (e.g., prolonged thawing of  $\text{Si(OH)}_4$  samples to allow depolymerization, three parallels measured). The median coefficient of variation among parallels was 5 % for  $[\text{NO}_3 + \text{NO}_2]$  and  $[\text{PO}_4]$ , 2 % for  $[\text{Si(OH)}_4]$ , and 9 % for  $[\text{NO}_2]$ ; higher coefficient of variation was observed when the absolute concentrations were low, e.g.,  $< 0.1 \mu\text{M}$ . Reproducibility was sufficient, and no parallels were excluded. Phosphate was analyzed, but N : P ratios for nutrients were, on average, 8 among all stations, suggesting that N was likely more important than P for potentially limiting primary production. These phosphate data ( $0.1\text{--}0.6 \mu\text{M}$  in the upper 50 m) are not discussed.

Samples for biogenic particulates and phytoplankton community composition were taken directly from the rosette and sediment traps. For bSiO<sub>2</sub> samples, 600 mL of seawater was collected from the rosette, filtered through a 1.2 µm polycarbonate filter (Millipore); for sediment trap material, less volume was necessary (e.g., 50–100 mL). Most bSiO<sub>2</sub> protocols use a 0.6 µm filter cutoff, e.g., Lalande et al. (2016); however, given the magnitude bSiO<sub>2</sub> quantified and the size range for regional diatoms we are confident that there was no meaningful systematic underestimate. After filtration, all samples were dried at 60 °C and stored until laboratory analysis using an alkaline digestion in Teflon tubes (Krause et al., 2009). For Chl *a*, water-column and sediment-trap samples were collected similarly, filtered on Whatman GF/F (0.7 µm pore size), and immediately frozen (−20 °C). In the laboratory, Chl *a* was extracted in 5 mL methanol in the dark at room temperature for 12 h. The solution was quantified using a Turner Designs 10 AU fluorometer, calibrated with Chl *a* standard (Sigma C6144), before and after adding two drops of 5 % HCl (Holm-Hansen and Riemann, 1978). Phytoplankton taxonomy and abundance samples were collected in 200 mL brown glass bottles from both the water column and sediment traps, immediately fixed with an aldehyde mixture of hexamethylenetetramine-buffered formaldehyde and glutaraldehyde at 0.1 % and 1 % final concentration, respectively, as suggested by Tsuji and Yanagita (1981), and stored cool (5 °C) and dark. Samples were analyzed with an inverted epifluorescence microscope (Nikon TE300 and Ti-S, Japan), using the Utermöhl (1958) method, in a service laboratory for diatom taxonomy (> 90 individual genera or species categories were identified) and abundance at the Institute of Oceanology Polish Academy of Science.

### 2.3 Rate measurements

Biogenic silica production was measured using the radioisotope tracer <sup>32</sup>Si. Approximately 150 or 300 mL samples, depending on the station biomass, were incubated with 260 Bq of high specific activity <sup>32</sup>Si(OH)<sub>4</sub> (> 20 kBq µmol Si<sup>−1</sup>). After addition, samples were transported to the onboard incubator and placed in neutral density screened bags, simulating 50 %, 20 %, and 1 % of irradiance just below the surface, for 24 h. After incubation, samples were processed immediately by filtering bottle contents through a 25 mm, 1.2 µm polycarbonate filter (Millipore) matching bSiO<sub>2</sub> filtrations. Each filter was then placed on a nylon planchette, covered with Mylar when completely dry, and secured using a nylon ring. Samples were aged into secular equilibrium between <sup>32</sup>Si and its daughter isotope, <sup>32</sup>P (~ 120 days). <sup>32</sup>Si activity was quantified on a GM multicounter (Risø National Laboratory, Technical University of Denmark) as described in Krause et al. (2011). A biomass-specific rate (i.e., *V*<sub>b</sub>) was determined by normalizing the gross rate (*ρ*) to the corresponding [bSiO<sub>2</sub>] at the same depth of collection using a logistic-growth approach (Kristiansen et al., 2000; Krause et

al., 2011). For bSiO<sub>2</sub> and *ρ*, values within a profile were integrated throughout the euphotic zone (i.e., surface to 1 % *I*<sub>0</sub>) using a trapezoidal scheme. A depth-weighted *V*<sub>b</sub> was calculated within the euphotic zone by integrating *V*<sub>b</sub> and dividing by depth-integrated values (Krause et al., 2013).

Two methods were used to assess whether ambient silicic acid (Si(OH)<sub>4</sub>) limited diatom Si uptake. The <sup>32</sup>Si activity additions, incubation conditions, and sample processing are as described above. At four stations (Edgeøya, Polar Front, Hinlopen, and Atlantic), eight 300 mL samples collected at a single depth within the euphotic zone were manipulated to make an eight-point concentration gradient between ambient and +18.0 µM [Si(OH)<sub>4</sub>]; the maximum concentration was assumed to saturate Si uptake. Si uptake has been shown to conform to a rectangular hyperbola described by the Michaelis–Menten equation:

$$V_b = \frac{V_{\max} [\text{Si(OH)}_4]}{K_S + [\text{Si(OH)}_4]}, \quad (1)$$

where *V*<sub>max</sub> is the maximum specific uptake rate and *K*<sub>S</sub> is half-saturation constant, i.e., concentration where *V*<sub>b</sub> = 0.5 × *V*<sub>max</sub>. Data were fit to Eq. (1) using a nonlinear curve fit algorithm (SigmaPlot 12.3). The second type of experiment used only two points: ambient and +18.0 µM [Si(OH)<sub>4</sub>]; four-depth profiles were done at three stations (Bellsund Hula, Hornsunddjupet, Erik Eriksenstretet). The ratio of Si uptake at +18.0 µM [Si(OH)<sub>4</sub>] to Si uptake at ambient [Si(OH)<sub>4</sub>] defines an enhancement (i.e., Enh) statistic. This two-point approach was conducted at all depths in the euphotic zone; Enh ratios > 1.08 imply kinetic limitation beyond analytical error given the methodology (Krause et al., 2012).

Net primary productivity (PP) was quantified concurrently with biogenic silica production at six stations at the depth of approximately 50 % of surface irradiance (Table 1). Carbon uptake rates were measured using a modification of the <sup>14</sup>C uptake method (Steeemann Nielsen, 1952). Water samples were spiked with 0.2 µCi mL<sup>−1</sup> of <sup>14</sup>C labeled sodium bicarbonate (Perkin Elmer, USA) and distributed in three clear plastic bottles and one dark (40 mL each). Subsequently, they were incubated for 24 h in the deck incubator with a 50 % light reduction mesh. After incubation, samples were filtered onto 0.2 µm nitrocellulose filters. The filters were stored frozen (−20 °C) in scintillation vials with 10 mL EcoLume scintillation liquid (MP Biomedicals LLC, USA) until further processing. Once on land, the particulate <sup>14</sup>C was determined using a scintillation counter (TriCarb 2900 TR, Perkin Elmer, USA). The carbon uptake values in the dark were subtracted from the mean of the triplicate carbon uptake values measured in the light incubations. Using contemporaneous *ρ* measurements and PP measurements, the diatom contribution to PP is estimated as follows:

$$\text{Diatom \% PP} = 100 \times \frac{\rho \times (\text{Si} : \text{C})^{-1}}{\text{PP}}, \quad (2)$$

**Table 1.** Station properties including surface temperature, nutrients and chlorophyll *a* ( $\pm$  standard deviation), 20 m integrated biogenic silica stock ( $\int$  bSiO<sub>2</sub>), production ( $f\rho$ ), depth-weighted specific production ( $V_{AVE}$ ), 40 m integrated diatom abundance ( $\int$  Diatom), and export of bSiO<sub>2</sub> and diatoms at 40 m. The disparity between the integration depths for bSiO<sub>2</sub> standing stock and diatom abundance reflects the lack of bSiO<sub>2</sub> samples to 40 m depth. Note: Hinlopen (ice) station not included. The Polar Front (ice) station is the mean of two profiles.

Station name	<i>T</i> (°C)	[NO <sub>3</sub> + NO <sub>2</sub> ] (μM)	[Si(OH) <sub>4</sub> ] (μM)	[Chl <i>a</i> ] (μg L <sup>-1</sup> )	20 m $\int$ bSiO <sub>2</sub> (mmol Si m <sup>-2</sup> )	20 m $f\rho$ (mmol Si m <sup>-2</sup> day <sup>-1</sup> )	20 m $V_{AVE}$ (day <sup>-1</sup> )	40 m $\int$ Diatom abundance (10 <sup>9</sup> cells m <sup>-2</sup> )	40 m bSiO <sub>2</sub> export (mmol Si m <sup>-2</sup> day <sup>-1</sup> )	40 m diatom export (10 <sup>6</sup> cells m <sup>-2</sup> day <sup>-1</sup> )
van Mijenfjorden <sup>a</sup>	-0.43	8.1	3.8	1.84 ± 0.19	10.8	—	—	7.67	9.03	769
Bredfjupet <sup>a</sup>	4.72	9.4	4.5	0.72 ± 0.03	1.9	0.27	0.13	—	—	—
Bellsund Hula	0.69	< 0.1	0.5	2.66 ± 0.05	15.3	0.49	0.06	—	—	—
Hornsund	-0.28	1.6	1.1	2.50 ± 0.20	—	—	—	8.97	—	1180
Hornsuddjupet <sup>a</sup>	-0.20	< 0.1	0.4	2.43 ± 0.17	42.2	1.46	0.03	—	—	—
Edgeøy <sup>a,c</sup>	-0.70	0	0.7	1.99 ± 0.03	—	—	—	—	—	—
Erik Eriksenstretet <sup>a</sup>	-1.58	0.4	0.4	4.77 ± 0.31	34.9	1.03	0.04	252	4.00	436
Polar Front station <sup>a,b</sup>	2.19	< 0.1	1.1	3.00 ± 0.03	—	—	—	527	—	—
Atlantic	4.10	3.3	1.4	6.66 ± 0.33	—	—	—	—	9.20	2380

<sup>a</sup> Denotes concurrent primary production and biogenic silica production measurements at one depth.

<sup>b</sup> 25 m depth.

<sup>c</sup> Surface values, no profile taken.

where the Si : C ratio for diatoms can be used from culture values. The most widely used Si : C ratio is 0.13 (Brzezinski, 1985); however, this study lacked polar diatom strains. Takeda (1998) grew two polar diatoms at 2 °C and in iron-replete media and reported Si : C from 0.10 to 0.18; however, this was extrapolated based on direct measurement of cellular N and converting using the Redfield–Ketchum–Richards C : N ratio of 6.6. A more recent study, Lomas and Krause (2018), reported data on 11 polar diatom species grown at 2 °C with direct measurement of biogenic silica and particulate organic carbon and nitrogen. For larger diatom species (> 1000 μm<sup>3</sup> biovolume) these authors observed the average Si : C was 0.25 ± 0.04 (standard error, SE), with a higher ratio for smaller species (< 1000 μm<sup>3</sup>) 0.32 ± 0.04 (SE). Most of the diatom assemblage during ARCEX was composed of larger cells; thus, we use Si : C of 0.25.

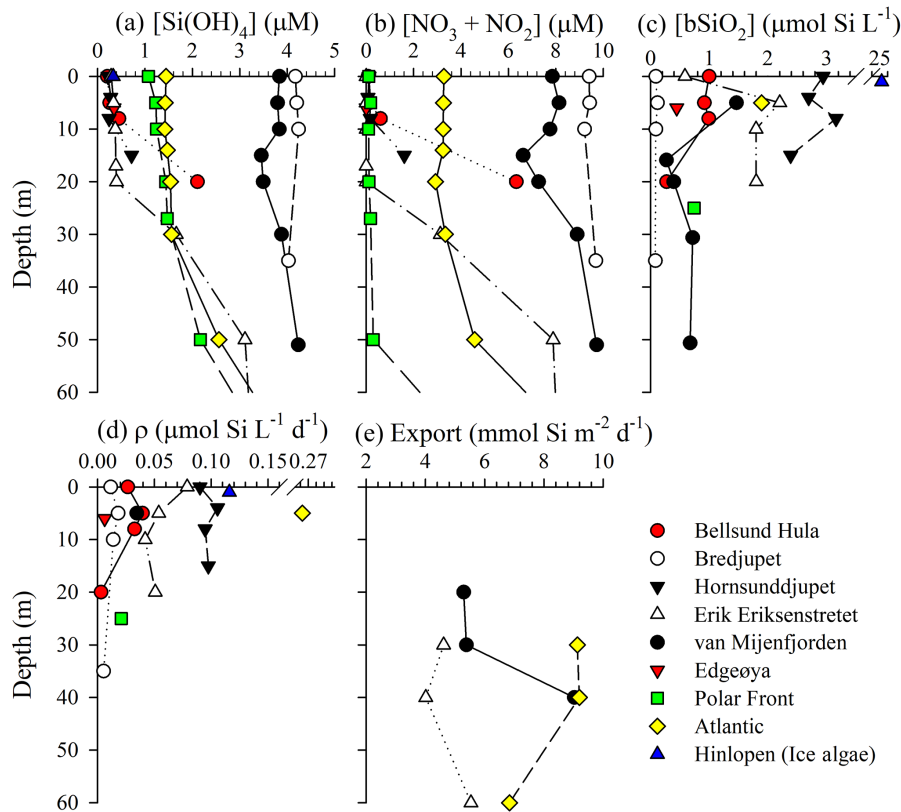
Export rates were calculated using the standing stock measurements, length of deployment, and trap opening area (0.004 m<sup>2</sup>). These approaches are common and detailed elsewhere (Wiedmann et al., 2014; Krause et al., 2009).

### 3 Results

#### 3.1 Hydrography and spatial patterns

The regional ecosystem around Svalbard is driven by ice dynamics (Sakshaug, 2004). A majority of the southern Svalbard archipelago had open water 1 week prior to the cruise open water, which was anomalous compared to similar dates in previous years (e.g., 2014, 2015, ice data archived at <http://polarview.met.no/>, last access: 29 October 2018). By the end of the cruise, Svalbard could have been entirely circled by the vessel, with only open drift ice in the northeastern region. While 2016 was among the lowest years for total Arctic sea ice, the ice extent in Svalbard and the Barents Sea is highly dynamic. Ice edges may be pushed southward into the Barents Sea proper by wind while areas to the north remain ice-free, e.g., Wassmann et al. (1999) and references therein.

Spatial patterns in hydrography and nutrients were highly variable. In the southwestern stations (e.g., fjords and Atlantic-influenced water), the surface temperature ranged between 1 and 4 °C; a similar temperature was observed in the Atlantic station south of the Polar Front (Fig. 1e). Northeastern domain stations were more influenced by Arctic water and the surface temperatures ranged between -2 and 1 °C (Fig. 1e). Surface nutrient concentrations, particularly [NO<sub>3</sub> + NO<sub>2</sub>] and [Si(OH)<sub>4</sub>], showed a broad range. The highest surface [NO<sub>3</sub> + NO<sub>2</sub>] was observed in the southwestern fjords, between 2 and > 8 μM, and the Atlantic station (~ 3 μM, Fig. 1a). The surface concentrations at the remaining stations were < 0.5 μM or near detection limits (Fig. 1a). [Si(OH)<sub>4</sub>] was lower than [NO<sub>3</sub> + NO<sub>2</sub>] (i.e., Si : N < 1) among stations where [NO<sub>3</sub> + NO<sub>2</sub>] was > 0.1 μM. At high [NO<sub>3</sub> + NO<sub>2</sub>] stations, the [Si(OH)<sub>4</sub>] ranged from 1.1–



**Figure 2.** Vertical profiles for (a) dissolved silicic acid, (b) nitrate + nitrite, (c) biogenic silica standing stock, (d) biogenic silica production rate, and (e) biogenic silica export. Symbols are associated by station, and line connectors are used to denote profile data opposed to individual symbols noting samples at one depth.

4.5  $\mu\text{M}$  (Fig. 1b) but the range was lower among other stations (0.4–1.1  $\mu\text{M}$ , Fig. 1b). The  $\text{bSiO}_2$  concentration (proxy for diatom biomass, Fig. 1c) was typically similar to, or lower than, surface  $[\text{Si}(\text{OH})_4]$ . The highest surface  $[\text{bSiO}_2]$  was observed in the southern stations (Atlantic-influenced waters):  $\sim 2\text{--}3 \mu\text{mol Si L}^{-1}$  (Fig. 1c). At most other stations the  $[\text{bSiO}_2]$  was  $< 1 \mu\text{mol Si L}^{-1}$ . Among all stations and depths  $\text{bSiO}_2$  varied by a factor of  $\sim 40$  (does not include Hinlopen ice algae).

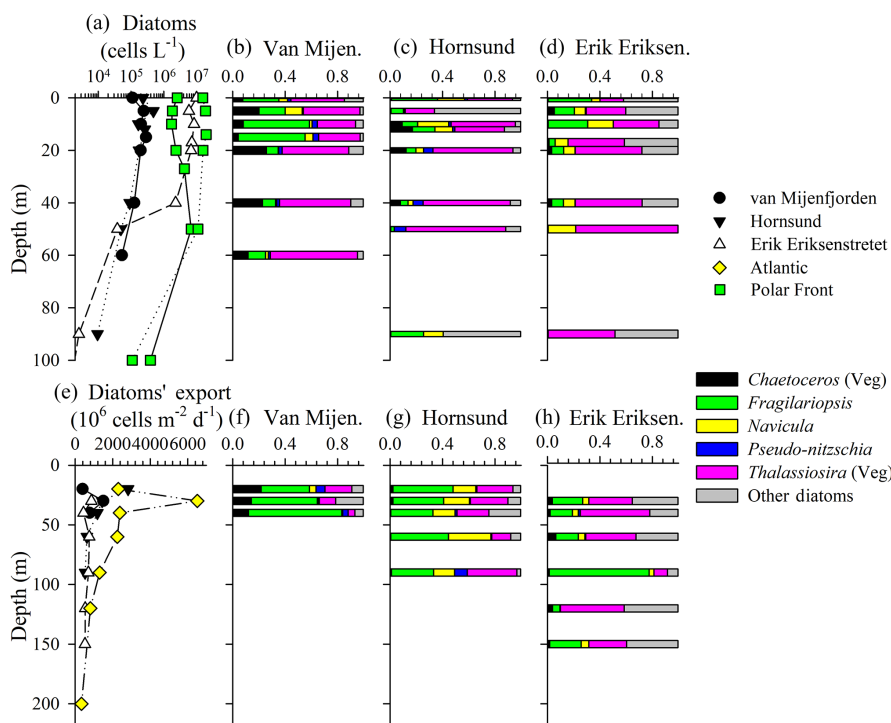
Primary productivity, measured at six stations at 5 m (approximately 50% of surface irradiance), varied by 2 orders of magnitude. The lowest primary productivity rates were observed at the four stations with the lowest surface  $[\text{NO}_3 + \text{NO}_2]$  and ranged from 2 to 13  $\mu\text{g C L}^{-1} \text{ day}^{-1}$ ; at these stations  $[\text{Chl } a]$  ranged from 2.0 to 4.8  $\mu\text{g L}^{-1}$  (Table 1, Fig. 1d). The highest rates were measured at van Mijenfjorden and Bredjupet:  $100 \pm 65$  and  $27 \pm 1 \mu\text{g C L}^{-1} \text{ day}^{-1}$ , respectively, and corresponded to high  $[\text{NO}_3 + \text{NO}_2]$  and low  $[\text{Chl } a]$  of 1.8 and 0.7  $\mu\text{g L}^{-1}$ , respectively (Table 1, Fig. 1d).

### 3.2 Vertical profiles

As expected, most stations showed strong vertical gradients in nutrient concentrations. Profiles in the southwestern re-

gion of Svalbard (van Mijenfjorden, Bredjupet) had elevated  $[\text{Si}(\text{OH})_4]$ , with little vertical structure. Vertical  $[\text{Si}(\text{OH})_4]$  profiles among other stations showed typical nutrient draw-down between the surface and  $\sim 20$  m. At these stations, surface  $[\text{Si}(\text{OH})_4]$  concentrations were typically  $< 1.5 \mu\text{M}$  and subsurface values (to 20 m) ranged from 0.5 to 3.0  $\mu\text{M}$  (Fig. 2a).  $[\text{NO}_3 + \text{NO}_2]$  exceeded  $[\text{Si}(\text{OH})_4]$  among all depths at five stations (Fig. 2b), whereas in the remaining stations  $[\text{NO}_3 + \text{NO}_2]$  exceeded  $[\text{Si}(\text{OH})_4]$  (i.e.,  $\text{Si} : \text{N} < 1$ ) at depths  $> 5$  m (Bellsund Hula),  $> 20$  m (Erik Erikstenstretet), and  $> 27$  m (Polar Front). For these latter three stations,  $[\text{NO}_3 + \text{NO}_2]$  had a significant drawdown in surface waters, but then increased with depth without a similar degree of vertical enhancement in  $[\text{Si}(\text{OH})_4]$  (Fig. 2).

The  $[\text{bSiO}_2]$  was typically highest at or near the surface, with a maximum of  $\sim 2 \mu\text{mol Si L}^{-1}$  (Fig. 2c). At the Bellsund Hula and Erik Erikstenstretet stations, subsurface  $[\text{bSiO}_2]$  maxima were present (Fig. 2c; note – no surface data are available for van Mijenfjorden). Among nonprofile stations,  $[\text{bSiO}_2]$  was within the range observed among vertical profiles except for the Hinlopen ice algae where ice, which was melted at ambient air temperature on the vessel, had exceptionally high  $[\text{bSiO}_2]$  (Fig. 2c). The integrated  $\text{bSiO}_2$  be-



**Figure 3.** Diatom abundance (a) and assemblage composition (b–d) in the water column, and diatom export (e) and assemblage composition (f–h) within sediment traps. Note – taxonomy information only shown for stations where both water-column and sediment-trap data were available (see text for species). Resting spores (e.g., *Chaetoceros*, *Thalassiosira*) were absent from the 40 m sediment traps; thus, proportional abundances for spore-producing taxa are entirely for vegetative cells. For (a), there are replicate diatom abundance measurements (from separate hydrocasts) for the Polar Front station.

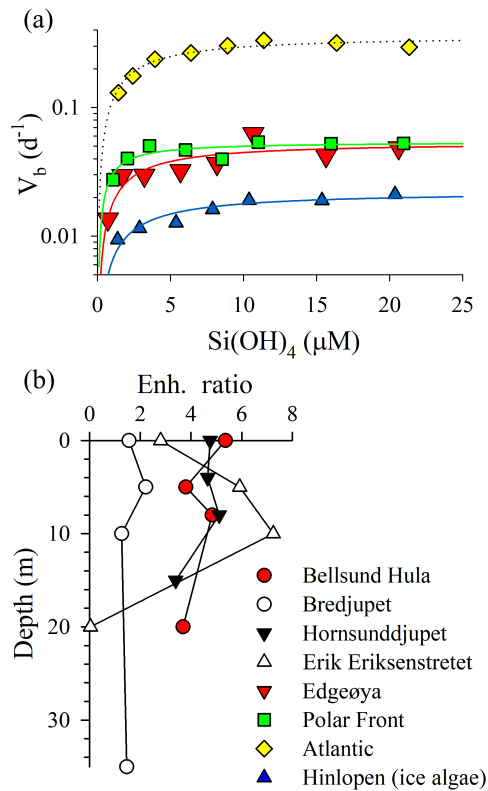
tween the surface and 20 m ( $\int \text{bSiO}_2$ ) spanned over an order of magnitude, with a low at Bredjupet ( $1.9 \text{ mmol Si m}^{-2}$ ) and a high at Hornsunddjupet ( $42.4 \text{ mmol Si m}^{-2}$ , Table 1) despite their proximity ( $\sim 50 \text{ km}$ ).

Diatom abundance and taxonomy data were sampled at fewer stations, but the vertical and spatial variability generally mirrored trends in  $[\text{bSiO}_2]$ . In the surface waters of van Mijenfjorden and Hornsund, diatom abundances ranged between  $5 \times 10^4$  and  $5 \times 10^5 \text{ cells L}^{-1}$  in the upper 50 m (Fig. 3a). However, within the same vertical layer at the Erik Eriksenstretet and Polar Front (duplicate profiles) stations, diatom abundances were enhanced by up to 2 orders of magnitude ( $4 \times 10^4$ – $4 \times 10^7 \text{ cells L}^{-1}$ , Fig. 3a). When integrated to 40 m depth ( $\int \text{Diatom}$ ), matching the shallowest sediment-trap depth among the three stations reported (Fig. 3e–h), diatom inventories also showed a variability of 2 orders of magnitude as observed in  $\int \text{bSiO}_2$ .  $\int \text{Diatom}$  was lowest at van Mijenfjorden ( $7.67 \times 10^9 \text{ cells m}^{-2}$ ) and highest at the Polar Front station ( $527 \times 10^9 \text{ cells m}^{-2}$ , Table 1).

Among the stations which had corresponding sediment trap deployments (van Mijenfjorden, Hornsund, Erik Eriksenstretet), the diatom-assemblage composition was similar despite differences in abundance. The van Mijenfjorden station was dominated by *Thalassiosira* (e.g., *T. antarctica*

*var. borealis*, *T. gravida*, *T. hyalina*, *T. nordenskiöldii*), *Fragilariopsis cylindrus*, and *Chaetoceros furcellatus* (Fig. 3b). *Chaetoceros* spp. were nearly absent from Erik Eriksenstretet (Fig. 3d) and of little importance at Hornsund (Fig. 3c). *Thalassiosira* species (same as van Mijenfjorden) cells also dominated Hornsund and Erik Eriksenstretet among most depths (Fig. 3c, d). However, at Hornsund, deeper depths were dominated by diatom groups less frequently observed (“Other diatom” category, Fig. 3), and with small contributions from *Fragilariopsis cylindrus* and *Navicula vanhoefenii*.

Diatom  $\text{bSiO}_2$  productivity,  $\rho$ , mirrored trends in biomass. Among the profiles, rates generally varied from  $\rho < 0.01$  to  $0.11 \mu\text{mol Si L}^{-1} \text{ day}^{-1}$  (Fig. 2d). The  $\rho$  was highest in the Atlantic station (Fig. 2d), which was expected given the higher  $\text{bSiO}_2$  (Fig. 2c). However, the rates in the Hinlopen ice algae were like those quantified at Hornsunddjupet,  $\sim 0.1 \mu\text{mol Si L}^{-1} \text{ day}^{-1}$ , despite the ice algae station having an order of magnitude more biomass. This suggests the Hinlopen ice algae were senescent or stressed and a sizable portion of the measured  $\text{bSiO}_2$  was nonactive or detrital. When integrated in the upper 20 m,  $\int \rho$  ranged from 0.27 to  $1.46 \text{ mmol Si m}^{-2} \text{ day}^{-1}$  (Table 1), which is a smaller proportional range than observed in  $\int \text{Diatoms}$  and  $\int \text{bSiO}_2$ . Over-



**Figure 4.** Assessment of Si uptake limitation by available silicic acid during ARCEX. **(a)** Eight-point kinetic experiments taken at four stations (legend next to **b**). Data were fit to a Michaelis–Menten hyperbola using SigmaPlot 12.3 software. **(b)** Enh. ratio profiles (i.e.,  $V_b$  in  $+18.0 \mu\text{M}$   $[\text{Si}(\text{OH})_4]$  treatment relative to  $V_b$  in the ambient  $[\text{Si}(\text{OH})_4]$  treatment) at four stations.

all,  $\text{bSiO}_2$ -normalized rates ( $V_b$ ) were low among all stations and depths ( $< 0.01$  to  $0.13 \text{ day}^{-1}$ ). The depth-weighted  $V_b$ , i.e.,  $V_{\text{AVE}}$ , had a narrower range of between  $0.03$  and  $0.13 \text{ day}^{-1}$ . Thus, doubling times for  $\text{bSiO}_2$  in the upper 20 m ranged between 5 and 23 days.

The rate of diatom biogenic silica production was kinetically limited by ambient  $[\text{Si}(\text{OH})_4]$  in 95 % of the samples examined. Full kinetic experiments verified that Si uptake conformed to Michaelis–Menten kinetics (Fig. 4a; adjusted  $R^2$  ranged  $0.64$ – $0.92$  among experiments). The highest  $V_{\text{max}}$  was observed in the Atlantic station ( $0.36 \pm 0.02 \text{ day}^{-1}$ ), which also had the highest ambient  $[\text{Si}(\text{OH})_4]$  among the full kinetic experiments ( $1.4 \mu\text{M}$ ).  $V_{\text{max}}$  observed at Edgeøya and the Polar Front was nearly identical ( $0.05 \pm < 0.01 \text{ day}^{-1}$  for both) and was lowest in the Hinlopen ice diatoms ( $0.02 \pm < 0.01 \text{ day}^{-1}$ ).  $K_S$  constants had a narrower range, with a low of  $0.8 \pm 0.3 \mu\text{M}$  at the Polar Front and between  $2.1$  and  $2.5 \mu\text{M}$  among the other three stations. Among these full-kinetic experiments, the Enh ratio ranged from  $1.8$  to  $7.7$ , with the most intense  $[\text{Si}(\text{OH})_4]$  limitation of uptake observed in the Hinlopen ice diatoms. For profiles where two-point kinetic ex-

periments were conducted, the same trends were observed (Fig. 4b). The Enh ratio was similar among depths at Bellsund Hula ( $1.5$ – $2.2$ ), Hornsunddjupet, and Bredjupet ( $3.4$ – $5.4$  for latter two stations, Fig. 4b). At Erik Erikssenstretet, Enh ratios were more variable, ranging from  $2.8$  to  $7.3$  in the upper 10 m, with no Enh effect (i.e.,  $< 1.08$ ) observed at 20 m – this was the only sample and depth which showed no resolvable degree of kinetic limitation for Si uptake.

Rates of  $\text{bSiO}_2$  and diatom export were variable. Among the three sediment trap regions,  $\text{bSiO}_2$  export rates ranged from  $\sim 4$  to  $10 \text{ mmol Si m}^{-2} \text{ day}^{-1}$  (Fig. 2e). These rates are significant and represent up to 50 % of the  $\int \text{bSiO}_2$  in the upper 20 m at van Mijenfjorden (Table 1). For diatom cells, a similar degree of variability was observed. Export at van Mijenfjorden ranged from  $390$  to  $1500 \times 10^6 \text{ cells m}^{-2} \text{ day}^{-1}$ , similar to ranges at Hornsund ( $520$ – $2800 \times 10^6 \text{ cells m}^{-2} \text{ day}^{-1}$ ) and Erik Erikssenstretet ( $510$ – $860 \times 10^6 \text{ cells m}^{-2} \text{ day}^{-1}$ , Fig. 3e). The Atlantic station had significantly higher diatom export ( $800$ – $2300 \times 10^6 \text{ cells m}^{-2} \text{ day}^{-1}$ ) among all depths in the upper 120 m (Fig. 3e). The  $\text{bSiO}_2$  and the export of diatom cells were highly correlated ( $r = 0.67$ ,  $p < 0.01$ ,  $n = 15$ ; Spearman’s rho test). Among all stations, *Fragilariopsis cylindrus* had the highest contribution to diatom export, and *Thalassiosira* species (e.g., *T. antarctica*, *T. gravida*, *T. hyalina*, *T. nordenskiöldii*) were also important (Fig. 3f–h). In Hornsund, *Navicula* (*N. vanhoefenii*, *N. sp.*) was an important genus for export (Fig. 3g), but this was not observed elsewhere. Similarly, “Other diatom” groups were proportionally important at Erik Erikssenstretet (Fig. 3h), as were *Thalassiosira* resting spores at the Atlantic station (data not shown). Among all diatoms, the only groups which were numerically important in both the water column and the sediment traps were *Fragilariopsis cylindrus* and *Thalassiosira* species (Fig. 3b–d, f–h).

## 4 Discussion

### 4.1 Diatom Si cycling relative to other systems

To our knowledge, this is the first report of  $\text{bSiO}_2$  production data of the natural diatom community in this sector of the Arctic. Other studies have reported  $\rho$  data in the subarctic Atlantic Ocean (Brown et al., 2003; Kristiansen et al., 2000; Allen et al., 2005)  $\sim 10$ – $20^\circ$  latitude south of our study region or in Baffin Bay (Hoppe et al., 2018; Tremblay et al., 2002). However, the Hoppe et al. (2018) study only includes  $\rho$  measured after a 24 h manipulation experiment and only at one site and depth near the Clyde River just east of Nunavut (Canada); no data are reported for the ambient conditions, and the measurements from Tremblay et al. (2002) are based on net changes in standing stocks instead of gross  $\text{bSiO}_2$  production. Banahan and Goering (1986) report the only  $\rho$  to date in the southeastern Bering Sea; however, Varela et al. (2013) recently reported that  $[\text{Si}(\text{OH})_4]$  in surface wa-



ters ( $> 5 \mu\text{M}$ ) is unlikely to be significantly limiting to diatoms in any sector of the Bering, Chukchi, or Beaufort Sea regions. Around Svalbard, some previous studies have examined other Si-cycling components, including variability in  $\text{bSiO}_2$  in the water column (Hodal et al., 2012) and sediments (Hulth et al., 1996),  $\text{bSiO}_2$  and diatom export (Lalande et al., 2016, 2013), or trends in  $[\text{Si}(\text{OH})_4]$  (Anderson and Dryssen, 1981). The  $\rho$  measurements presented here have no straightforward study for comparison; therefore, we compare these to the previous high-latitude Atlantic data and to well-studied sectors of the Southern Ocean.

During our study,  $\int \rho$  in the Svalbard vicinity was low. In the northeastern Atlantic between Iceland and Scotland, the reported  $\int \rho$  ranged between 6 and  $166 \text{ mmol Si m}^{-2} \text{ day}^{-1}$  (Brown et al., 2003; Allen et al., 2005). These rates are significantly higher than at our four profile stations (Table 1), and the degree of difference does not appear to be driven by differences in integration depth (compared to our study, Table 1). Given the higher  $[\text{Si}(\text{OH})_4]$  in the southern region of the Atlantic subpolar gyre (Hátún et al., 2017), the maximum achievable  $\int \rho$  may vary with latitude. While our profile sampling was opportunistic, it appears we sampled some stations with significant diatom biomass (high  $\int \text{bSiO}_2$ ), but the corresponding production rates ( $\int \rho$ ) were low, with estimated doubling times on the order of 11–23 days. This suggests these high-biomass stations may have been near, or past, peak bloom conditions (Fig. 2a, b) and the seasonal timing is consistent with regional field and modeling studies inferring diatom bloom dynamics from Chl *a* trends (Wassmann et al., 2010; Oziel et al., 2017). Kristiansen et al. (2000) reported  $\rho$  in Oslofjorden during the late winter (February–March) ranging from 0.03 to  $2.0 \mu\text{mol Si L}^{-1} \text{ day}^{-1}$ , over nine sampling periods with corresponding  $V_b$  of between  $< 0.01$  and  $0.28 \text{ day}^{-1}$ ; however, this system has a higher  $\text{Si}(\text{OH})_4$  supply and surface concentration at the start of the bloom period ( $> 6 \mu\text{M}$ ), approximately 50 % higher than the highest surface concentrations observed during our study (Fig. 2a). The specific rates observed in our study fall within the lower values reported by Kristiansen et al. (2000), which may be explained by the reduced uptake from lower  $[\text{Si}(\text{OH})_4]$  (e.g., Fig. 4).

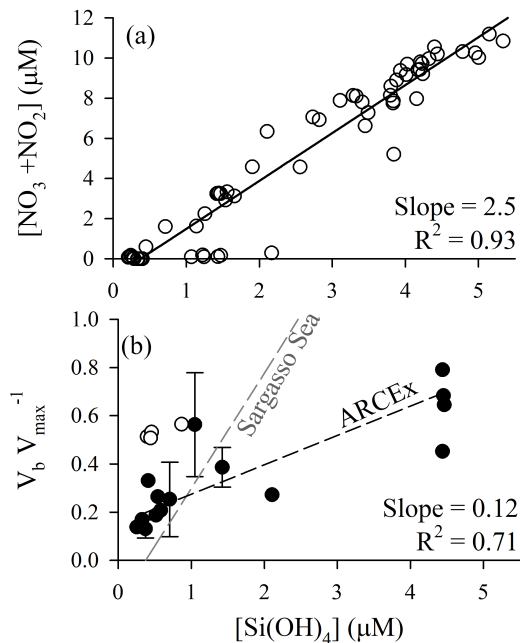
The Southern Ocean is one of the most globally significant regions for production of  $\text{bSiO}_2$ . The surface  $[\text{Si}(\text{OH})_4]$  and  $[\text{NO}_3 + \text{NO}_2]$  are among the highest in the ocean and the source waters usually have  $> 50 \%$  excess  $\text{Si}(\text{OH})_4$  relative to nitrate (Brzezinski et al., 2002). Thus, exceptional  $\text{Si}(\text{OH})_4$  drawdown relative to nitrate is required for diatom biomass yield to be limited by Si in this region. The mean  $\int \rho$  in sectors of the Southern Ocean are variable. In the Weddell Sea, winter rates range between 2.0 and  $3.2 \text{ mmol Si m}^{-2} \text{ day}^{-1}$  in the seasonal ice zone (Leynaert et al., 1993). Within the sub-Antarctic zone, rates averaged 1.1 and  $4.8 \text{ mmol Si m}^{-2} \text{ day}^{-1}$  in the summer and spring, respectively (Fripiat et al., 2011). At the terminus of diatom blooms in the sub-Antarctic and polar frontal zone, rates

can be lower, e.g.,  $0.1\text{--}0.3 \text{ mmol Si m}^{-2} \text{ day}^{-1}$  (Fripiat et al., 2011); such values are similar to the range observed during our study, especially since these Southern Ocean studies integrated  $\int \rho$  deeper than 40 m (e.g., 50–100 m). Brzezinski et al. (2001) reported average  $\int \rho$  of  $\sim 25 \text{ mmol Si m}^{-2} \text{ day}^{-1}$  (integrated from surface to 80–120 m) during intense blooms in the seasonal ice zone, which propagated south of the Antarctic polar front. But despite the massive diatom  $\text{bSiO}_2$  accumulating in these blooms,  $V_{\text{AVE}}$  generally ranged between 0.05 and  $0.15 \text{ day}^{-1}$  (Brzezinski et al., 2001). Given the order-of-magnitude difference in  $[\text{Si}(\text{OH})_4]$  and  $\int \rho$  between the Arctic and Southern Ocean, the similar  $V_{\text{AVE}}$  in both regions may be more reflective of thermal effects on diatom growth rate, since Si uptake and diatom growth rates are tightly coupled, or of a significant accumulation of detrital  $\text{bSiO}_2$  (i.e., diatom fragments) in the Southern Ocean, where low temperatures reduce  $\text{bSiO}_2$  remineralization rates (Bidle et al., 2002).

#### 4.2 Potential for silicon limitation of diatom productivity

Suboptimal silicon availability affects the rate of diatom  $\text{bSiO}_2$  production and can limit their growth. For diatoms in Svalbard and the broader region of the subpolar and polar European Atlantic, both  $[\text{Si}(\text{OH})_4]$  and its availability relative to N appear to be suboptimal for creating intense diatom blooms, such as those occurring in the Southern Ocean. Yet, the Arctic spring bloom is consistently dominated by diatoms or *Phaeocystis* (Degerlund and Eilertsen, 2010), which suggests some level of adaptation for diatoms to the low  $[\text{Si}(\text{OH})_4]$  environment. Stoichiometry of silicon availability relative to nitrate can help diagnose Si limitation; the most widely accepted diatom Si : N ratio is  $\sim 1$  based on temperate and low-latitude clones (Brzezinski, 1985). The average Si : N ratio for two polar diatom clones (silicic acid and iron replete) reported in Takeda (1998) was  $0.96 \pm 0.24$  (SE). A more recent culture study by Lomas and Krause (2018), reported Si : N for 11 polar diatom clones grown at  $2^\circ\text{C}$  among exponential and stationary growth phases, in both replete and N-limiting nutrient conditions; these authors observed Si : N among all clones, treatments, and nutrient conditions ( $> 150$  data points) was  $1.7 \pm 0.10$  (SE).

The silicon kinetic data provide clarity for interpreting Si and N nutrient drawdown. Diatoms have an *r*-selected ecological strategy and are typically the first phytoplankton group to bloom in this region under stratified conditions (Reigstad et al., 2002). The 1.7 Si : N from Lomas and Krause (2018) for nutrient-replete polar diatoms suggests they consume 70 % more Si relative to N. However, under kinetic limitation, diatoms have long been inferred to reduce Si per cell in culture to avoid growth limitation (Paasche, 1973) – this was recently observed directly in the field for the first time (McNair et al., 2018). Given the clear kinetic limitation observed during ARCEX (Fig. 4), this likely reduced the di-



**Figure 5.** Diagnosis of potential silicon limitation for diatom production during ARCEX. **(a)** Nitrate + nitrite drawdown as a function of dissolved silicic acid. **(b)** The ratio of  $V_b$  at ambient  $[\text{Si}(\text{OH})_4]$  to  $V_{\text{max}}$  versus dissolved silicic acid. In both panels, linear regressions were done using a model II reduced major axis method; for panel **(b)** the regression line does not include the Hornsunddjupet station (open circles). For comparison, the same relationship for the Sargasso Sea in the North Atlantic subtropical gyre is shown, as synthesized in Krause et al. (2012).

atom Si : N ratio to closer to the canonical 1 : 1 ratio. Thus, the kinetic limitation in this region may result in N and Si being consumed in near-equal amounts (i.e., Si : N  $\sim$  1) and previous inferences of diatom processes based on 1 : 1 Si : N drawdown appear valid.

Nutrient relationships support the potential for silicon to be a controlling factor of regional diatom productivity. When plotting  $[\text{NO}_3 + \text{NO}_2]$  as a function of  $[\text{Si}(\text{OH})_4]$  (Fig. 5a) a few trends emerge. (1) The slope of the linear regression relationship ( $2.5 \pm 0.1 \text{ mol N} (\text{mol Si})^{-1}$ ) denotes that  $\text{NO}_3 + \text{NO}_2$  is consumed at over twice the rate per unit  $\text{Si}(\text{OH})_4$ . (2) Given that the source water  $[\text{NO}_3 + \text{NO}_2]$  concentration is only roughly twice that of  $[\text{Si}(\text{OH})_4]$ , a 2.5 drawdown ratio would predict  $\text{NO}_3 + \text{NO}_2$  to be depleted before  $\text{Si}(\text{OH})_4$ . The latter observation suggests that  $\text{Si}(\text{OH})_4$  could be the yield-limiting nutrient for diatoms during a spring bloom period only if they dominate the phytoplankton assemblage and consume Si : N in ratios  $> 1$ , e.g., 1.7 as reported by Lomas and Krause (2018). Field data demonstrate interannual variability. Nitrate and silicic acid drawdown within the upper 50 m during the spring season (1980–1984) was discussed by Rey et al. (1987) who suggested apparent nitrate limitation (1980, 1981) and silicic acid limitation (1983,

1984). The Reigstad et al. (2002) analysis of nitrate and silicic acid drawdown in the central Barents Sea shows similarities to ARCEX in that the diatom assemblage could only draw down  $[\text{Si}(\text{OH})_4]$  to  $\sim 1 \mu\text{M}$  (May 1998) and  $\sim 0.5 \mu\text{M}$  (July 1999). These authors suggest that physical effects on phytoplankton explain the variability, where diatoms dominate in shallow mixed waters as opposed to *Phaeocystis pouchetii* dominating in deeper mixed waters. Clearly, interannual and local differences in mixing, which may favor *Phaeocystis pouchetii* over diatoms (Reigstad et al., 2002), can affect the assemblage and nutrient drawdown trajectory (e.g., see points with high  $[\text{Si}(\text{OH})_4]$  and  $[\text{NO}_3 + \text{NO}_2]$  close to detection limit, Fig. 5a); therefore, diagnosis of whether Si limits diatom production should be accompanied by additional analyses.

When considering the European sector of the Arctic and sub-Arctic between 60 and 80° N, there is compelling evidence that ambient  $[\text{Si}(\text{OH})_4]$  limits the rate of diatom  $\text{bSiO}_2$  production. During ARCEX, the relationship between  $V_b$  and  $[\text{Si}(\text{OH})_4]$  supports that Si regulates diatom productivity to some degree (Fig. 4). Our kinetic data demonstrate that in three of four experiments  $K_S$  was  $\sim 2.0 \mu\text{M}$ , but in the Polar Front the  $K_S$  was lower  $\sim 0.8 \mu\text{M}$ . These data are consistent with community kinetic experiments reported in Oslofjorden where  $K_S$  and  $V_{\text{max}}$  were between 1.7 and 11.5  $\mu\text{M}$  and between 0.16 and 0.64  $\text{day}^{-1}$ , respectively, with the lowest  $V_{\text{max}}$  observed during the declining diatom bloom (Kristiansen et al., 2000). These authors concluded that silicon ultimately controlled diatom productivity during this bloom (Kristiansen et al., 2001). In the only other kinetic experiments reported in the northeast Atlantic, Brown et al. (2003) and Allen et al. (2005) observed linear responses in  $V_b$  between ambient and 5  $\mu\text{M}$   $[\text{Si}(\text{OH})_4]$ , which suggests uptake did not show any degree of saturation at this concentration (Note: the single experiment reported in Allen et al. (2005) is one of four experiments originally reported in Brown et al., 2003). These field-based  $K_S$  values are considerably higher than parameters used in Barents Sea models, e.g., 0.5  $\mu\text{M}$  (Slagstad and Støle-Hansen, 1991) and 0.05  $\mu\text{M}$  (Wassmann et al., 2006), which reflect the high-efficiency Si uptake reported for cultures (Paasche, 1975). Fitting a regression to the  $V_b V_{\text{max}}^{-1}$  as a function of  $[\text{Si}(\text{OH})_4]$  (line shown in Fig. 5b) suggests that 2.8  $\mu\text{M}$  is the best constrained half-saturation concentration (i.e., concentration where  $V_b V_{\text{max}}^{-1} = 0.5$ ) for the regional assemblage. This empirical value excludes the Hornsunddjupet assemblage (white symbols, Fig. 5b), and their inclusion decreases this aggregated half-saturation to 2.3  $\mu\text{M}$ . Unlike diatoms in the North Atlantic Subtropical Gyre, e.g., the Sargasso Sea (Krause et al., 2012), regional diatoms do not appear to be well adapted for maintaining  $V_b V_{\text{max}}^{-1} > 0.5$  at low  $[\text{Si}(\text{OH})_4]$ . Instead, diatoms during the spring season appear to be best adapted for concentrations exceeding 2.3  $\mu\text{M}$ . It is plausible that as  $[\text{Si}(\text{OH})_4]$  is depleted, diatoms may slow growth from severe limitation of Si

uptake (Fig. 5b) and/or biomass yield (i.e., stock of diatom  $b\text{SiO}_2$  far exceeds  $\text{Si}(\text{OH})_4$ ).

To avoid growth limitation under conditions of kinetic limitation (i.e., suboptimal  $[\text{Si}(\text{OH})_4]$ ), diatoms can reduce their silicon per cell. A guideline from culture work is that diatoms can alter their silicon per cell by a factor of 4 (Martin-Jézéquel et al., 2000). Thus, when uptake is reduced to  $< 25\%$  of  $V_{\text{max}}$  (i.e., concentration which promotes uptake at half the half-saturation level) diatoms must slow growth to take up enough Si to produce a new cell. Using the empirical half-saturation constant range ( $2.3\text{--}2.8\ \mu\text{M}$ ), calculated from Fig. 5b and using Eq. (1) to solve for the concentrations where  $V_b/V_{\text{max}}^{-1} \leq 0.25$  ( $V_{\text{max}}$  is a constant), suggests that when  $[\text{Si}(\text{OH})_4]$  is below  $0.3\text{--}0.8\ \mu\text{M}$ , the degree of kinetic limitation could force diatoms to slow growth in response. This type of limitation could occur even if diatom  $b\text{SiO}_2$  stock was not sufficiently high to induce yield limitation, e.g., it could not deplete all  $\text{Si}(\text{OH})_4$  from the assemblage undergoing one division. Such a range is lower than the common interpretation of the Egge and Aksnes (1992) data set showing diatoms may be outcompeted by flagellates when  $[\text{Si}(\text{OH})_4] < 2\ \mu\text{M}$ , a value which is more reflective of an ecological niche opposed to a physiological threshold, as has been purported in numerous citations of these data. At these inferred limiting  $[\text{Si}(\text{OH})_4]$  there would be up to  $0.8\ \mu\text{M}$   $[\text{NO}_3 + \text{NO}_2]$  remaining (Fig. 5a), which would allow nonsiliceous phytoplankton to draw down the remaining N. Therefore, under shallow stratified conditions which favor diatoms over *Phaeocystis* (sensu Reigstad et al., 2002),  $[\text{Si}(\text{OH})_4]$  may regulate regional diatom productivity through either yield- or severe-kinetic limitation. This provides the most direct assessment to date, supporting the general ideas proposed for Si regulation of regional diatom productivity (Rey, 2012; Rey et al., 1987; Reigstad et al., 2002).

### 4.3 Diatom contribution to primary production

Among the six sites with paired PP and  $\rho$  measurements, the bloom phase can be inferred from the magnitude of nutrient drawdown,  $[\text{Chl } a]$ , PP, and  $p\text{CO}_2$  (data not shown). Bredjupet appeared to be a pre-bloom station given the high surface nutrient concentrations, while the van Mijenfjorden station appeared to be in an early bloom phase based on relative high nutrients and moderate  $[\text{Chl } a]$ . The Erik Eriksenstretet station represented a peak bloom condition, whereas assemblages at Hornsunddjupet and Edgeøya appeared to be post-bloom and in a stage of decline. The Polar Front station represented the end or late-phase bloom condition; however, at this station *Phaeocystis* was abundant (data not shown), suggesting it may have dominated the bloom dynamics instead of diatoms.

The diatom contribution to PP was highly variable. Among the stations with high  $[\text{NO}_3 + \text{NO}_2]$  (van Mijenfjorden, Bredjupet) the diatom contribution to PP (e.g., Eq. 2) was low, at  $2\text{--}3\%$ . At two stations, Hornsunddjupet and the Polar

Front, the diatom contribution to PP increased to  $25\text{--}30\%$ . In the Edgeøya and Erik Eriksenstretet stations, diatoms accounted for a majority or all of PP, at  $70\%$  and  $180\%$ , respectively. Given that diatoms can reduce their cellular Si in response to kinetic limitation (Paasche, 1973; McNair et al., 2018), the Si : C ratio of 0.25 based on nutrient-replete polar diatoms in culture may systematically underestimate diatom contribution to PP using our approach. For example, if kinetic limitation reduced Si per cell by  $50\%$  (i.e.,  $V_b/V_{\text{max}}^{-1} \approx 0.50$ , Fig. 5b) but did not affect cellular C, then the Si : C ratio would be 0.13 (i.e., temperate Si : C diatom value), and nearly all the calculated diatom contributions would double. Considering the degree of kinetic limitation at most stations (Fig. 5b), this suggests our estimates are conservative except at Erik Eriksenstretet. The unrealistic value at Erik Eriksenstretet underscores the issue with the Si : C ratio (Eq. 2); however, adjusting Si : C downward would increase the diatom contribution. Thus, there may have been lower C per cell for diatoms at this station due to other factors associated with the phase of the bloom and/or the different assemblage; e.g., *Porosira glacialis* was more abundant at this station and has a large vacuole which could lower C content, thereby increasing Si : C (data not shown).

Clearly, diatoms can play a significant role in local productivity, but these data demonstrate a “boom and bust” nature. At stations at or near peak bloom levels (e.g., Edgeøya, Erik Eriksenstretet), diatoms could account for nearly all primary production. However, they may also contribute an insignificant percentage of primary production prior to the onset of the bloom (e.g., van Mijenfjorden, Bredjupet). But even when physical conditions may favor *Phaeocystis* blooms, diatoms appear to be significant contributors to primary production (Polar Front station). In such a situation, N would be predicted to be the limiting nutrient as it will be consumed by both *Phaeocystis* and diatoms whereas Si will only be consumed by the latter.

In the European Arctic, shifts in summer-period phytoplankton communities away from diatom-dominated conditions have been observed in numerous studies. One of the most important observations has been the increasing abundances of *Phaeocystis* in ice-edge (Lasternas and Agustí, 2010) or under-ice blooms (Assmy et al., 2017). These changes have corresponded with larger-scale shifts in the export of diatoms to depth in the Fram Strait (Nöthig et al., 2015; Lalande et al., 2013; Bauerfeind et al., 2009). The timing of these shifts, e.g., mid-2000s, correspond with the broader regional reduction in winter mixed-layer  $[\text{Si}(\text{OH})_4]$  concurrent with the shift to negative gyre-index state in the latter half of the decade (Hátún et al., 2017). With a reduction in pre-bloom  $\text{Si}(\text{OH})_4$  supply, diatoms may run into limitation sooner during the bloom cycle and thus leave more residual nitrate for nondiatom phytoplankton. Given the highly variable contribution of diatoms to primary productivity, resolving a climate change or natural physical oscillation signal will be challenging. A similar conclusion about detecting

a climate-change signal was made in the eastern Bering Sea by Lomas et al. (2012) given the natural variability in primary production.

#### 4.4 Diatoms and export

The bSiO<sub>2</sub> export rates observed during ARCEX were significant relative to the standing stocks. At van Mijenfjorden, the rate of export in the upper 40 m represented 39 % of the ∫ bSiO<sub>2</sub> standing stock (23.3 mmol Si m<sup>-2</sup>, integral of data in Fig. 2c) in the same vertical layer. This quantity was much higher than at Erik Eriksenstretet, where the 40 m export rate was < 11 % of the ∫ bSiO<sub>2</sub> in the upper water column (note: no samples were taken deeper than 20 m; thus, additional bSiO<sub>2</sub> between 20 and 40 m would lower the 11 % estimate). Given that the van Mijenfjorden site was located within shallow fjord waters (bottom depth approximately 60 m), such a high proportion export relative to standing stock may suggest either lateral focusing processes (e.g., discussed by DeMaster, 2002) and/or resuspension of sediment bSiO<sub>2</sub> into the water and resettlement. The rate of bSiO<sub>2</sub> export among all export and production stations was also at least a factor of 4 higher than ∫ ρ in the upper 20 m (Table 1). It is likely that some fraction of ∫ ρ was missed due to lack of sampling between 20 and 40 m, but with a less light at these depths, it is unlikely systematic underestimates of ρ caused the disparity. Given the deeper water at the Erik Eriksenstretet and Atlantic stations, such high bSiO<sub>2</sub> export may be driven by previously high ρ and bSiO<sub>2</sub> standing stock that accumulated in the overlying waters or, given the dynamic circulation in the region, this signal may have been laterally advected to these station locations.

Relative to previous studies, the bSiO<sub>2</sub> export rates were also high. During May 2012 in Kongsfjorden, Lalande et al. (2016) reported bSiO<sub>2</sub> export rates between 0.2 and 1.3 mmol Si m<sup>-2</sup> day<sup>-1</sup> in the upper 100 m. A similar range was observed by Lalande et al. (2013) in the eastern Fram Strait using moored sediment traps (2002–2008) collecting at depths between 180 and 280 m. Lalande et al. (2013) concluded that, despite warm anomaly conditions, pulses of bSiO<sub>2</sub> export were positively correlated to the presence of ice in the overlying waters, which stratifies the water and helps initiate a diatom bloom. However, if the light was insufficient to stimulate a bloom, Lalande et al. (2013) suggested much of the pulse of bSiO<sub>2</sub> exported to depth may have originated in the ice and sank during melting. Indeed, the low V<sub>b</sub> (< 0.01 day<sup>-1</sup>) observed at the Hinlopen station (ice algae), despite the moderate ρ measured (0.12 μmol Si L<sup>-1</sup> day<sup>-1</sup>), suggests that most of the ice-associated bSiO<sub>2</sub> was detrital and not associated with living diatoms. Thus, the recent ice retreat observed prior to the ARCEX cruise was a potential source of such high bSiO<sub>2</sub> export to depth despite the considerably lower ∫ ρ in the upper 20 m.

Among the groups examined, the most important diatom genera for standing stock and export were *Thalassiosira* and

*Fragilariopsis*, suggesting these groups are important drivers of bulk bSiO<sub>2</sub> fluxes. Given the large-size and chain-forming life histories for the dominant species within each genus, it is likely that their dominance in the trap abundances helps explain the high correlation ( $r = 0.67$ ,  $p < 0.01$ ; Spearman's rho test) between bSiO<sub>2</sub> and diatom export. Given this degree of correlation, it would be expected that both bSiO<sub>2</sub> and diatom export would be similarly enhanced relative to previous studies; however, this was not observed.

Comparing the magnitude of bulk bSiO<sub>2</sub> export and the export of diatom cells suggests significant food web repackaging occurred. The export of diatom cells in Kongsfjorden (Lalande et al., 2016) were similar to or a factor of 3 lower than rates quantified during ARCEX (Table 1, Fig. 3e), whereas bSiO<sub>2</sub> export during ARCEX was over an order of magnitude higher than bSiO<sub>2</sub> export in Kongsfjorden. One possible explanation for the higher degree of bSiO<sub>2</sub> export enhancement, relative to diatom-cell export, between studies is that more exported material during ARCEX was repackaged and modified in the food web. For instance, in Erik Eriksenstretet gel traps confirm the presence of aggregates and mesozooplankton fecal pellets (Ingrid Wiedmann, data not shown), and in van Mijenfjorden detrital particles and sediment material were most prominent on the gel traps opposed to clearly recognizable material (e.g., diatom valves). This repackaging is consistent with previous observation in the Barents Sea showing high potential for copepod fecal pellets to be exported in the Polar Front and Arctic-influenced regions during spring (Wexels Riser et al., 2002). It also supports the general ideas for the importance of diatom organic matter in fueling secondary production regionally during this season (Degerlund and Eilertsen, 2010, and references therein).

## 5 Conclusion

This is the first regional data set with contemporaneous measurements of diatom bSiO<sub>2</sub> standing stock, production, and export and assessment of kinetic limitation by [Si(OH)<sub>4</sub>] in the European Arctic. Among stations and depths there was widespread limitation of diatom bSiO<sub>2</sub> production rates by ambient [Si(OH)<sub>4</sub>] during spring-bloom conditions. The kinetic parameters for diatom Si uptake (e.g., K<sub>S</sub>) quantified in our study are significantly higher than rates used in regional models and quantified in polar diatom cultures; therefore, these data will help future modeling efforts better simulate diatom–Si dynamics. Given the trajectories of Si and N consumption, diatom-dominated blooms (vs. *Phaeocystis*-dominated) could deplete Si(OH)<sub>4</sub> prior to nitrate (yield limitation); and at some stations, the degree of kinetic limitation by ambient [Si(OH)<sub>4</sub>] could have resulted in diatom growth being slowed. Diatom contribution to PP was highly variable, ranging from < 10 % to ~ 100 % depending on the bloom stage; but even when *Phaeocystis* appeared to be favored,

diatoms still had a significant ( $\sim 25\%$ ) contribution to PP. While there was agreement with previous regional studies regarding the export rate of diatom cells, we observed significantly elevated bSiO<sub>2</sub> export. Such a discrepancy can be resolved if a higher fraction of the diatom material exported during our study was modified by zooplankton grazers, relative to previous studies, or if much of this bSiO<sub>2</sub> was derived from melting ice and/or advection.

*Data availability.* All data are available upon request to the authors or are available through the UiT research data bank (<https://dataverse.no/dataverse/uit>, last access: 29 October 2018).

*Author contributions.* JWK, CMD, and SA conceived and designed the study and conducted analysis. JWK, CMD, IAM, PA, MFM, IW, and SA conducted the fieldwork. PW and SK conducted analysis. All coauthors contributed to the writing of the paper, led by JWK.

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

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