



Synechococcus in the Atlantic Gateway to the Arctic Ocean

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Increasing temperatures, with pronounced effects at high latitudes, have raised questions about potential changes in species composition, as well as possible increased importance of small-celled phytoplankton in marine systems. In this study, we mapped out one of the smallest and globally most widespread primary producers, the picocyanobacterium Synechococcus, within the Atlantic inflow to the Arctic Ocean. In contrast to the general understanding that Synechococcus is almost absent in polar oceans due to low temperatures, we encountered high abundances (up to 21,000 cells mL^{-1}) at 79°N, and documented their presence as far north as 82.5°N. Covering an annual cycle in 2014, we found that during autumn and winter, Synechococcus was often more abundant than picoeukaryotes, which usually dominate the picophytoplankton communities in the Arctic. Synechococcus community composition shifted from a quite high genetic diversity during the spring bloom to a clear dominance of two specific operational taxonomic units (OTUs) in autumn and winter. We observed abundances higher than 1000 cells mL⁻¹ in water colder than 2°C at seven distinct stations and size-fractionation experiments demonstrated a net growth of Synechococcus at 2°C in the absence of nano-sized grazers at certain periods of the year. Phylogenetic analysis of petB sequences demonstrated that these high latitude Synechococcus group within the previously described cold-adapted clades I and IV, but also contributed to unveil novel genetic diversity, especially within clade I.

Keywords: picocyanobacteria, picoeukaryotes, temperature adaptation, *petB* sequences, flow cytometry, high latitude ecosystems, Svalbard, West Spitsbergen Current

INTRODUCTION

The widely abundant picocyanobacterium *Synechococcus* is estimated to be responsible for about 17% of ocean net primary productivity and thus to have a high impact on ocean ecosystems and biogeochemical cycles (Flombaum et al., 2013). *Synechococcus* is normally not considered to be bloom-forming even though they can appear in abundances as high as $1.2-3.7 \times 10^6$ cells mL⁻¹ in the Costa Rica dome (Saito et al., 2005). Using 37,699 discrete global *Synechococcus* observations between 69°S and 81°N and quantitative niche models, Flombaum et al. (2013) demonstrated temperature to be the main environmental parameter explaining the global distribution of *Synechococcus*. Accordingly, the regional range of temperature was found to be a relatively good predictor for the seasonal change in *Synechococcus* abundance (Tsai et al., 2013). Although the

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marine *Synechococcus* that have been studied in culture so far have a temperature optimum ranging from 20 to 33° C depending on the clade (Pittera et al., 2014), the highest annual average *in situ* cell abundances were found at temperatures around 10° C in the Indian and western Pacific Oceans, with averages of 34,000 and 40,000 cells mL⁻¹, respectively (Flombaum et al., 2013).

There has been observations of *Synechococcus* at low temperatures e.g., $<4^{\circ}$ C in low numbers (<100 cells mL⁻¹; Waterbury et al., 1986) and $<2^{\circ}$ C (Shapiro and Haugen, 1988; Gradinger and Lenz, 1995), but they are still often considered to be nearly absent from the polar ocean (Pedrós-Alió et al., 2015) in contrast to cold adapted eukaryotic picophytoplankton that occur in high abundances both in Arctic (Sherr et al., 2003; Lovejoy et al., 2007; Tremblay et al., 2009; Zhang et al., 2015) and in Antarctic waters (Doolittle et al., 2008). Only a few studies have actually documented *Synechococcus* north of 70° and none have so far described the genetic diversity of these northern populations or tested their temperature optimum.

During four expeditions Gradinger and Lenz (1995) observed maximal abundances of 5500 Synechococcus cells mL^{-1} in the Atlantic inflow to the Arctic Ocean west of Svalbard at 78°N, while they did not find any Synechococcus cells in surface samples of polar water (defined as water having Temp $< 0^{\circ}$ C; Salinity < 34). Further south, following a transect from 70.5 to 74°N, Not et al. (2005) recorded a maximum abundance of 25,000 cells mL⁻¹ in the Norwegian and Barents Seas in August 2002. In the western Canadian Arctic, Cottrell and Kirchman (2012) found abundances of 40-80 cells mL⁻¹ in coastal waters of the Chukchi Sea and the Beaufort Sea at 71.5°N, both during summer and winter cruises. Nelson et al. (2014) concluded in their overview that the Synechococcus distribution in this region is controlled mainly by inflow of the relatively warm Pacific water, but argue that water temperature alone cannot be used to define environments in which Synechococcus may reside as they do persist at water temperatures near the freezing point $(-1.8^{\circ}C)$ (Nelson et al., 2014).

Synechococcus is often found in Arctic lakes and rivers, and freshwater runoff may thus also represent a source of Synechococcus cells to the Arctic Ocean (Vincent et al., 2000). Using 16S rRNA analysis, Waleron et al. (2007) revealed that picocyanobacteria present in the Canadian Beaufort Sea originate from the Mackenzie River and other nearby inflows. High abundances of *Synechococcus* (30,000 cells mL^{-1}) were also found in the Laptev Sea, but were restricted to brackish waters near the Lena River delta, while further away from the delta, abundances decreased with increasing salinity to a total absence at salinities >20 (Moreira-Turcq and Martin, 1998). All these studies support Waterbury et al. (1986) claiming that only few brackish species tolerate wide salinity ranges and that many strains are obligate marine. Assuming that Atlantic Synechococcus have a low tolerance to salinity changes, the question remains whether the low salinity in the Arctic surface waters constrains their distribution in the polar ocean.

The Atlantic inflow is the main conveyor, not only of water and heat, but also of more southern species into the Arctic Ocean. *Synechococcus* has accordingly been suggested as a bio-indicator for the advection of Atlantic waters into the Arctic Ocean



colored according to the month of sampling: January (black), March (white), May (green), August (orange), and November (blue). Star-symbols indicate sampling of water for fractionated growth experiments. Hexagon symbols with white border indicate sampling for molecular analysis. Red arrows indicate the main flow of the West Spitsbergen Current drawn after Cokelet et al. (2008), Randelhoff et al. (submitted).

(Murphy and Haugen, 1985; Gradinger and Lenz, 1995). The main transport follows the West Spitsbergen Current (WSC) which is an extension of the Norwegian Atlantic Current splitting up into two branches around 79-80°N (Figure 1). The WSC is about 100 km wide and is confined over the continental slope along the Norwegian coast. It has an average speed of 10 cm s⁻¹ (Cokelet et al., 2008) but can reach a speed of up to 24- 35 cm s^{-1} (Boyd and D'Asaro, 1994; Fahrbach et al., 2001). The inflow follows a strong annual cycle with maximum volume transport during winter (20 Sv in February) and minimum during summer (5 Sv in August, Fahrbach et al., 2001) (N.B. the unit sverdrup (Sv) is equal to 1 million $m^3 s^{-1}$). Strong variations in the strength of the Atlantic inflow combined with varying sea ice extension make it challenging to assess the spread of Atlantic organisms in this area. Little is known about how Synechococcus populations, originating from the Norwegian coast or further south, are affected as they are transported into the Arctic Ocean or whether some Synechococcus lineages are favored under the transition to more Arctic conditions.

Temperature is one of the main drivers of *Synechococcus* biogeography. Among the five globally dominating *Synechococcus* lineages (clades I, II, III, IV, and CRD1), clades I and IV dominate at high latitudes in cold and coastal waters, while clades II and III are mostly found in warm, (sub)tropical areas (Zwirglmaier et al., 2008; Farrant et al., 2016; Sohm et al., 2016). Populations adapted to distinct thermal niches were also identified within the CRD1 clade, including one co-occurring with clade I and IV in cold, mixed waters of the Pacific Ocean (Farrant et al., 2016).

Increasing ocean temperature in high latitude systems has drawn attention towards the growth of invasive organisms with higher temperature optima and subsequent ecosystem changes. In marine systems, small phytoplankton are expected to become relatively more abundant with warming (Morán et al., 2010; Tremblay et al., 2012) and it has been speculated that the warming of the Arctic Ocean could lead to a shift from picoeukaryotes to picocyanobacteria, with implications for food quality (Vincent, 2010). Flombaum et al. (2013) projected up to a 50% increase in Synechococcus at 60°N by the end of the twenty first century. Their models were however not able to make projections for higher latitude systems because observations in these areas are scarce. The aim of the present study is therefore to examine the distribution of Synechococcus in relation to environmental parameters and other microbial plankton groups within the Atlantic gateway to the Arctic. The genetic diversity of Synechococcus populations was also unveiled using a high resolution genetic marker, the *petB* gene (encoding the cytochrome b_6 subunit), in order to trace the geographical origin and seasonal changes of these populations.

MATERIALS AND METHODS

Locality and Sampling

This study covers the eastern part of the Fram Strait, where Atlantic water (AW) is transported northward by the West Spitsbergen Current (WSC). Data were collected during five cruises in 2014: January (06.01-15.01), March (05.03-10.03), May (15.05-02.06), August (07.08-18.08), and November (03.11-10.11). Transects were made across the core of AW inflow at 79 and 79.4°N during May, August and November. Further north (80.5 to 82.6°N) we investigated the WSC southern branch into the Arctic Ocean in January, March and August (Figure 1). The choice of sampling area and stations was largely determined by the extension of the sea ice (Figure 3). Vertical profiles of temperature, salinity and fluorescence were recorded on each sampling occasion using a SBE 911plus system. Water masses were defined based on the criteria presented in Table 1. Discrete water samples for analyses of nutrients (NO $_2^-$ + NO_3^- , NH_4^+ , PO_4^{3-} , H_4SiO_4) and enumeration of phytoplankton, viruses, bacteria, and heterotrophic nanoflagellates (HNF) were collected from 11 depths (1, 5, 10, 20, 30, 50, 100, 200, 500, 750, and 1000 m) using 10 L Niskin bottles. During the summer cruises we collected additional samples from the Deep Chlorophyll Maximum (DCM) (when different from any of the standard depths). The shallow shelf stations were sampled to near bottom and with higher sampling resolution in the surface.

Flow Cytometry

Abundances of pico- and nano-sized phytoplankton, viruses, bacteria and HNF were determined on an Attune[®] Acoustic Focusing Flow Cytometer (Applied Biosystems by Life technologies) with a syringe-based fluidic system and a 20 mW 488 nm (blue) laser. Samples were fixed with glutaraldehyde

TABLE 1 | Criteria determining the water masses.

| Water masses | Temperature (°C)/or density (kg m $^{-3}$) | Salinity |
|---------------------------|---|----------|
| Atlantic water (AW) | >2°C | >34.9 |
| cold Atlantic water (cAW) | 0 <t<2°c< td=""><td>>34.9</td></t<2°c<> | >34.9 |
| Intermediate water (IW) | <0°C | >34.9 |
| Arctic water (ArW) | $>27.7 \text{kg m}^{-3}$ | <34.9 |
| Surface water (SW) | $<\!27.7{\rm kg}{\rm m}^{-3}$ | <34.9 |
| Polar water (PW) | <0°C | <34.7 |

For further explanation see Cokelet et al. (2008). PW overlaps with SW and ArW.

(0.5% final conc.) at 4°C for a minimum of 2 h, flash frozen in liquid nitrogen and stored at -80°C until analysis, except in November, when phytoplankton was enumerated using fresh samples. For analysis of HNF the samples were stained with SYBR Green I (Molecular Probes, Eugene, Oregon, USA) for 2h in the dark and a minimum of 1 mL was measured at a flow rate of 500 μ l min⁻¹ following the protocol of Zubkov et al. (2007). The HNF population was discriminated from nano-sized phytoplankton based on green vs. red fluorescence and from large bacteria on a plot of side scatter vs. green fluorescence following the recommendations of Christaki et al. (2011). Pico- and nano-sized phytoplankton were counted directly after thawing and the various groups discriminated based on their side scatter signals (SSC) vs. orange fluorescence (Figure 2A; Marie et al., 1997; Larsen et al., 2001) as well as their red vs. orange fluorescence (Figure 2B). Synechococcus was identified in plots of orange fluorescence vs. side scatter signals (Figure 2A). For samples with low abundance of phytoplankton (March and January) a volume of 1.5 mL was counted, while 0.5 mL was sufficient for May, August and November-samples. Regular blank measurements using Milli-Q[®] water were made to ensure that there was no carry over of cells between samples and that electronic noise did not disturb the counts. Due to the inherent uncertainty connected to enumeration of cells when concentrations are low, we only included samples with > 20 cells mL^{-1} when relating counts to other environmental parameters (Figure 5). Samples for which 0-20 cells were detected (i.e., mainly those deeper than 500 m) are included in our total data set (Table S1, Figure S2).

Microscopy

The presence of *Synechococcus* was also confirmed by fluorescence microscopy (**Figure 2C**). Samples were fixed and stored as for flow cytometry. The samples were thawed, filtered onto Anodisc filters (Whatman, pore size $0.2 \ \mu$ m) and stained with SYBR Green I (Molecular Probes Inc., Eugene, Oregon) according to Patel et al. (2007). The samples were viewed and photographed at 400X using a Zeiss Axio Imager Z1 microscope with AxioCam MRm BW-camera, extended focus, epifluorescence illumination (HXP Illuminator) and Zeiss filter sets 09 and 43 for SYBR Green and chlorophyll fluorescence, respectively.



Size-Fractionated Growth Experiments

Water fractionation experiments were used to examine interaction between different size groups of microorganisms and to estimate growth rates of the different microbial components (Simek and Chrzanowski, 1992; Jürgens et al., 2000; Christaki et al., 2001; Sato et al., 2007). Experiments were performed once every cruise using water collected from 20 m (in August and May this depth was near DCM) at stations on the shelf (marked on **Figure 1**). The water was gently screened through 3, 5, 10, and 90 μ m mesh size filters by reverse filtration in order to successively exclude grazers of different sizes and thus create communities with increasing "top-predators" sizes. Water from each filtration treatment was gently transferred into triplicate 3.9 L transparent polycarbonate bottles (Nalgene[®]) by staggered filling using silicone tubing. The incubation experiments ran for 5 to 10 days but we show data only from the initial 5 days of incubation in order to better represent dynamics of the initial communities. Incubation water was sampled daily for enumeration of microorganisms and every second day for nutrients. Prior to setup, all bottles, carboys and silicon tubs were acid washed and then rinsed with Milli-Q® water. During the summer cruises (May and August) the experimental bottles were incubated on deck in plexiglass tanks with seawater flow-through (continuously pumped from 7 m depth), keeping the temperature close to in situ (May: 1.7 \pm 1.6°C and August: 1 \pm 0.8°C). A nylon net was wrapped around each bottle to reduce the PAR to about 30% of the surface irradiation. In the winter months (January, March, November), incubations were kept in a cooling room at a constant temperature of 2°C and in darkness, except in March were an in situ light cycle was set (16h darkness and 8h at 5 μ mol photons m⁻² s⁻¹). The fractionation experiments provided net growth rates of *Synechococcus* and HNF by fitting exponential functions to the change in the abundance of cells every 24 h during the first 5 days of the experiments (**Figure S1**).

DNA Extraction, PCR Cloning and Phylogenetic Analysis

Environmental samples for molecular analysis were collected by filtering water onto 0.22 μ m pore size Millipore[®] Sterivex filters. The filters were immediately flash frozen in liquid nitrogen and stored at -80°C until extraction. DNA and RNA were extracted simultaneously using the AllPrep DNA/RNA Mini Kit (Qiagen, Hilden, Germany) according to manufacturer's instructions with some optimisation for extraction from Sterivex filters as follows. The filters were thawed on ice and 1 mL extraction buffer (990 μ l RLT buffer; containing guanidine isothiocyanate + 10 μ L ßmercaptoethanol) was added before incubating for 4 min on a Vortex adapter at medium speed. The resulting lysates were recovered using a 10 mL syringe and used for nucleic acids extraction. DNA samples harvested from Arctic surface water collected in May (80°N, 10.7°E at 1 m depth, 10 L water filtered), August (80°N, 10.8°E at 1 m, 7.5 L filtered) and November (79°N, 6°E, 20 m, 20 L filtered) were selected to amplify the Synechococcus petB marker gene (stations marked on Figure 1 and profiles of picophytoplankton are included in Figure S2). Polymerase chain reaction (PCR) were performed using the petB primers and set-up recommended by Mazard et al. (2012) using 30-40 amplification cycles (iCycler, Bio-Rad, CA, USA). Positive PCR products were purified using the Zymo DNA Clean and ConcentratorTM-5 kit (Zymo research, CA, USA) and subsequently cloned with the StrataCloneTM PCR Cloning Kit (Agilent Technologies, CA, USA) following the manufacturer's instructions. A total of 96 clones from each of the three samples were picked (total 288 clones) and sequenced by LCG Genomic GmbH (Berlin, Germany) using Sanger sequencing. A total of 229 *petB* sequences were obtained and deposited in the GenBank database (accession no. KX345947-KX346174). These sequences are in the following referred to as "MicroPolar sequences."

The 229 MicroPolar sequences include 174 unique fulllength sequences. Together with 721 petB sequences from a non-redundant reference database (representing most of the genetic diversity so far identified within Prochlorococcus and Synechococcus genera; Farrant et al., 2016), the MicroPolar sequences were used to define operational taxonomical units (OTUs) at 97% identity using Mothur v1.34.4 (Table S2). Since all MicroPolar sequences clustered with clades I and IV reference sequences, a subset of the *petB* database, comprising only the 117 reference sequences of these clades, as well as the 174 unique MicroPolar petB sequences was used for a subsequent analysis. Phylogenetic reconstructions were based on multiple alignments of *petB* nucleotide sequences generated using MAFFT v7.164b with default parameters (Katoh and Standley, 2014). A maximum likelihood tree was inferred using PHYML v3.0, (Guindon and Gascuel, 2003) with the HKY + G substitution model, as determined using jModeltest v2.1.4 (Darriba et al., 2012) and estimation of the gamma distribution parameter of the substitution rates among sites and of the proportion of invariable sites. The tree was drawn using iTOL (Letunic and Bork, 2007). The 229 sequences retrieved from MicroPolar were recruited using BLASTN (v2.2.28+) against the full *petB* database: reads with more than 90% of their sequence aligned and with more than 80% sequence identity to their BLASTN best-hit were taxonomically assigned to their best-hit and subsequently used to build per-sequence read counts tables. Counts were then aggregated by OTUs and relative abundance was computed for each MicroPolar station.

Nutrients

Unfiltered seawater was filled directly from the Niskin bottles into 30 mL acid washed HDPE bottles and stored at -20° C. Nitrite and nitrate (NO₂⁻ + NO₃⁻), phosphate (PO₄³⁻) and silicic acid (H₄SiO₄) were measured on a Smartchem200 (by AMS Alliance) autoanalyser following procedures as outlined in Wood et al. (1967) for NO₃⁻ + NO₂⁻, Murphy and Riley (1962) for PO₄³⁻ and Koroleff (1983) for the determination of H₄SiO₄. The determination of NO₃⁻ was done by reduction to NO₂⁻ on a builtin cadmium column, which was loaded prior to every sample run. Seven-point standard curves were made prior to every run. Two internal standards and one blank were inserted for every 8 samples and these were used to correct for any drift in the measurements. Concentration of NH₄⁺ was determined directly in fresh samples using ortho-phthaladehyde according to Holmes et al. (1999).

RESULTS

Synechococcus cells were detected by flow cytometry in all samples within the upper 100 m of the water column during all seasons (**Figure 3** and **Figure S2**). The identity of *Synechococcus* was confirmed by epifluorescence microscopy (**Figure 2C**) and by sequencing of the *petB* gene (**Figure 7**). The closely related genus *Prochlorococcus* was never detected.

Synechococcus Distribution

The highest sampling frequency was obtained in May and August, when the sampling sites were restricted to latitudes below 80°N (May) and 81°N (August) by sea ice (**Figure 3**), while the most northern samples were acquired in January and March at around 82.5°N. *Synechococcus* was present in abundances higher than 50 cells mL⁻¹ in 337 samples both within the Atlantic water, Arctic water, Surface water and Polar water (water mass definitions are shown in **Table 1**). Within the cold surface water (<2°C, upper 50 m), 60% of the samples contained *Synechococcus* was not detected in the cold Atlantic water or intermediate water masses, which comprise water collected deeper than 500 m (see temperature-salinity plots in **Figure 3**).

In January, the average abundance in the upper 100 m was 51 cells mL^{-1} , with highest abundance found at 100 m depth (maximum 106 cells mL^{-1}) and generally low numbers in the surface (**Figures 3F**, **4A**). The lowest average abundance of



FIGURE 3 | (Left panel) lce-maps provided from the Norwegian Meteorological Institute (istjenesten@met.no) from following dates; January 10, March 7, May 23, August 12, and November 7, 2014. Transects shown in the middle panel as contour plots are marked with black boxes. (Middle panel) Contour plots showing the abundance (cells mL^{-1}) of *Synechococcus* and salinity of the upper 200 m from 4 cruises. January transect stretches from North-South between 15 and 20°E, while the remaining transects expands West-East (2–11°E) following the 79°N latitude isoline (see transect marked in boxes left of plots). The horizontal light blue lines above the plots roughly mark the cover of open drift ice. Note different scales for *Synechococcus* abundance. (Right panel) Potential temperature and salinity (TS) diagram for each month. Data included for all depths 1–1000 m. *Synechococcus* abundance is given on the z-axis by color gradient (N.B. different scales). Potential density (σ_0 , kg/m³) isolines overlaid with gray and the surface freezing line is show in dashed blue. Following water masses (Table 1) are marked: Atlantic water (AW), cold Atlantic water or Intermediate water (cAW or IW), which consist mainly of deep water samples (>500 m), Arctic water (ArW), surface water (SW) and Polar water (PW).



Synechococcus was found in March with 13 cells mL⁻¹ and a maximum abundance that did not exceed 40 cells mL^{-1} (Figures 3H, 4B). In May the maximum abundance was around 1300 cells mL⁻¹ and the average \pm SD was 181 \pm 147 cells mL⁻¹ (n = 150; Figures 3J, 4C). The highest *Synechococcus* abundances were detected in August with a maximum of 21,300 cells mL^{-1} . When averaged for the upper 50 m at most southern stations $(79-79.4^{\circ}N)$, abundances were 5700 ± 4200 cells mL⁻¹ (n = 61) (Figure 3L) over the whole transect, while abundances at the stations north of 80°N averaged to about 3000 \pm 2000 cells mL^{-1} (n = 27). In November Synechococcus cells were evenly distributed down to 200 m (Figure 4E), with a maximum of 1000 cells mL⁻¹ and an average abundance of 600 ± 250 cells mL⁻¹ (n = 18) within the upper 200 m. The vertical distribution of Synechococcus varied from mainly surface peaks in May (upper 20 m) to maximum abundance at depths greater than 50 m in August and November, to a more vertically uniform distribution in January and March with maxima in abundance at around 100 m depth (Figures 4A–E and Figure S2).

Biotic and Abiotic Environment

The association between phytoplankton abundances and environmental parameters showed that the abundance of both Synechococcus and picoeukaryotes decreased with increasing latitude, but that picoeukaryotes were relatively more abundant at the northernmost stations (Figure 5A). No clear relationship was found for salinity (ranging from 31 to 35 in this study), although the highest Synechococcus abundances were found at salinities >34.5, while picoeukaryotes had their peak abundance at lower salinities of 33.5-34 (Figure 5B). Further, we found picoeukaryotes to be strongly dominant over Synechococcus in 14 out of 17 samples with lowest salinity (31-33), all sampled in August. The abundance of Synechococcus ranged from 250 to 4000 cells mL^{-1} in these low salinity samples (Figure 3S). The presence of sea ice had no clear effect on the vertical distribution of picophytoplankton but at the ice-covered stations, a subsurface maximum of Synechococcus was most prominent. On the other hand, picoeukaryotes tended to peak near surface in ice-covered stations in March and May, while in August the highest surface



FIGURE 5 | (Two upper panels) Data collected within upper 100 m from all cruises plotted to see how abundance of *Synechococcus* (red) relates to environmental parameters, such as (A) latitude (B) salinity (C) temperature (D) chlorophyll fluorescence and (E–G) concentration of different sources of mineral nitrogen. Additionally (A,B,E–G) contain abundance of picoeukaryotes (green) and (D) nanophytoplankton (blue). Notice log-scale for (A,B,E–G). (Lower panel) (H–K) *Synechococcus* abundance (mL⁻¹) plotted against HNF abundance during the different months (indicated by color) within the upper 200 m. The broken line illustrates the one-to-one line. Notice different y-axis between months. Only samples with *Synechococcus* abundance >20 cells mL⁻¹ were included.

maximum of picoeukaryotes was found within the freshwater lens at stations without ice-cover (Figure S2).

The highest water temperatures were measured in August followed by those measured in January. The lowest surface temperatures were recorded in the ice-influenced surface waters in March and May. Temperature was the only parameter that displayed a strong relationship with *Synechococcus* abundance resulting in an exponential fit ($r^2 = 0.66$, p < 0.005, n = 346; Figure 5C), while picoeukaryotes did not show a similar strong relationship ($r^2 = 0.31$, p < 0.005, n = 372; Figure 5C). *Synechococcus* was more dominant at stations with low chlorophyll *a* (i.e., chl *a* fluorescence) compared to larger nanophytoplankton, which correlated positively to chl *a* (Figure 5D).

Nutrients were evenly distributed over the upper 200 m in the winter months (January to November), although a slightly lower concentration was observed in March within the upper 100 m (**Figures 4F,G,J**). In May and August all nutrients were depleted in the upper 10–20 m, with NO_3^- reaching the lowest values (**Figures 4H,I**). NH_4^+ reached

the highest values around 2 μM in August at depths below 20 m. At high N concentrations (>2 μM ; Figure 5E) Synechococcus were generally less abundant than picoeukaryotes, while under low N conditions they were equally numerous. When looking at the N sources separately it appears that at NH_4^+ > 0.5 μM , Synechococcus increased at higher NH_4^+ levels, whereas they decreased with increasing NO_3^- + NO_2^- (Figures 5F,G).

The abundance of HNF increased during the summer months, from less than 200 cells mL^{-1} in the winter months up to 1000 and 1500 HNF mL^{-1} in May and August, respectively (**Figures 5H–K**). *Synechococcus* and HNF abundances generally showed a positive relationship within the upper 100 m. In January and March, *Synechococcus* and HNF cell numbers were within the same order of magnitude, but with slightly more HNF than *Synechococcus* (i.e., below the dotted line; **Figure 5H**). In May highest *Synechococcus* abundances were found at the lowest HNF abundance and vice versa (**Figure 5I**). In August, *Synechococcus* was generally 10 times more abundant than HNF, a trend also observed in November, although less pronounced (**Figures 5J,K**).

Growth and Microbial Interactions

Net-growth rates of Synechococcus and HNF were estimated from four different size fractions (<3, <5, <10, and $<90 \ \mu$ m) from each of the five cruises and summarized in Figure 6 (for abundances during incubation see Figure S1). Synechococcus showed positive net growth in 9 out of 20 experiments mainly in January and March. Positive growth rates ranged from 0.01 to 0.13 d⁻¹. HNF showed positive growth in 14 out of 20 experiments, displaying a maximum growth rate of 0.45 d^{-1} when water was filtered through a 5 μ m mesh (Treat < 5 μ m) in January, otherwise the highest HNF growth rates were measured in May ranging from 0.13 to 0.3 d^{-1} . In January, November, and August HNF growth was reduced to close to zero after filtering in the Treat $< 90 \ \mu m$ and in March HNF showed negative growth in all treatments. Synechococcus showed positive growth in January, March in all size-fractions and in the Treat < 90 μm in May, which became strongly dominated by *Phaeocystis* sp. and where both HNF, picoeukaryotes and heterotrophic bacteria increased in abundance simultaneously. Synechococcus had the strongest negative growth in August and in the Treat $< 3 \ \mu m$ in May (Figure 6). In summary, we measured a positive growth of Synechococcus and negative growth of HNF in March, but in general negative Synechococcus and positive HNF growth in May, August, and November. Only in January and in the May <90 µm treatment, both Synechococcus and HNF displayed positive net growth. Corresponding to the seasonal changes in abundance (Figures 5H-K) the prey:HNF ratio (prey being the sum of all picoplankton; Synechococcus + picoeukaryotes + heterotrophic bacteria) of the initial community was highest in May i.e., most prey per HNF grazer and lowest in August, when HNF were more abundant. Generally, the maximum growth rates of Synechococcus were found when prey:HNF was at its highest (Figure S1).

Synechococcus Diversity

The gene *petB*, which has proved to display a high taxonomic resolution for picocyanobacteria (Mazard et al., 2012), was used as phylogenetic marker for Synechococcus genetic diversity. Only petB sequences related to clade I and IV were retrieved from our dataset (MicroPolar). Based on a petB reference database (including 117 sequences from clade I and IV, described in Farrant et al., 2016), enriched with the 174 unique petB sequences retrieved from MicroPolar samples, 41 OTUs were defined at 97% ID within clade I and IV (Figure 7). The petB sequences obtained in the present study correspond more specifically to sub-clades Ib and IVb, with a clear dominance of subclade Ib. Although none of these 41 OTUs form a new subclade, 17 OTUs were composed of only MicroPolar sequences and were not represented in the previous reference database (colored branches, Figure 7). In May sub-clade Ib was the only one present, whereas subclade IVb appeared in August and increased in relative abundance in November, indicating seasonal changes in the community composition. Seasonality was also found within subclade Ib. The majority of sequences obtained in August and November belonged to two specific OTUs (Arctic732-2b_Ib_IA and Arctic732-35b_Ib_IA), which mostly gather reference sequences from the Barents Sea ("Arctic,"



FIGURE 6 | Net growth rates (d⁻¹) of *Synechococcus* plotted against HNF net-growth. Net growth rates are obtained from the fractionation experiments from each cruise where an exponential growth curve was fitted to the change in abundance of the respective cells during a 5-day period (Figure S1). The color indicates the month and the legend at each point indicates the size-fraction treatment from which the values were obtained.

72.5°N, 19.57°W) and the North Atlantic Ocean (The Extended Ellett Line; "EEL," 57–63°N). In contrast, sequences retrieved from samples harvested in May were more evenly distributed over all other OTUs defined within subclade Ib that mainly gather sequences from The Atlantic Meridional Transect ("AMT," http://www.amt-uk.org/) and the North Sea ("MICROVIR" cruise 50–60°N).

In order to assess whether the genetic populations sampled in MicroPolar cruises could be related to other cold-water populations, we also recruited Illumina reads from 62 surface water metagenomes collected during the Tara Oceans cross-ocean ecosystem study using the same petB database (Karsenti et al., 2011; Farrant et al., 2016) (https://doi.pangaea. de/10.1594/PANGAEA.840718, note that the Tara Oceans samples analyzed here do not include recent Arctic samples from the latest Tara Ocean cruise as they are not yet published). These data showed that the two most abundant MicroPolar OTUs in subclade Ib (Arctic732-2b_1b_1A and Arctic732-35b_1b_1A) had a low relative abundance in Tara Oceans stations. Other OTUs identified in MicroPolar samples were also poorly represented in the Tara Oceans dataset, with the notable exception of subclade Ib OTUs, "MP may P1 1m E08 Ib I" two "MP_may_P1_1m_D10_Ib_IA" (formed and only of MicroPolar sequences), that were dominant in Tara Oceans coldest stations (<14°C), and of the subclade "Ellet21_IVb_IVC" present in the IVb OTU Tara Oceans dataset at all temperatures and especially at cold ($<14^{\circ}$ C) and intermediate (18–22°C) temperatures. Other MicroPolar OTUs were detected at a similarly low level in all temperature ranges of Tara Oceans stations.



DISCUSSION

Arctic Adaptation; Synechococcus vs. Micromonas

For the first time we here documented a high abundance of Synechococcus in the Atlantic gateway to the Arctic Ocean north of 79°N. Synechococcus is generally not thought to be part of the picophytoplankton community in Arctic water masses (e.g., Pedrós-Alió et al., 2015), which has repeatedly been found to be dominated by picoeukaryotes, such as Micromonas spp. (Not et al., 2005). Li et al. (2013) do document their existence in the Canadian Basin of the Arctic proper although as a very small fraction (2%) of the total picophytoplankton community at the only one station higher north than 70°. Arctic Micromonas spp. differ from Micromonas genotypes identified elsewhere in the World Ocean (Lovejoy et al., 2007), with these Arctic types being adapted to low temperatures. Similarly, by combining our observations with data from the Tara Ocean we confirmed the latitudinal shift previously described within the Synechococcus genus between the warm-adapted clades II and III and the coldadapted clades I and IV (Zwirglmaier et al., 2008; Mazard et al., 2012; Farrant et al., 2016). Interestingly, clade IV was clearly dominating in Atlantic waters from the Tara Oceans dataset and its relative contribution seemed to increase with temperature in August and November in MicroPolar samples, while clade I appeared to dominate in colder Arctic waters. Thus, although this would need to be confirmed by physiological characterization of representative strains, it suggests that clade I could be adapted to colder waters than clade IV. Overall, it seems that temperature is the main driver of *Synechococcus* abundance and diversity in this area.

In laboratory experiments using isolates from tropical sites, Synechococcus has been found not to grow at temperatures below 10°C (Mackey et al., 2013), even though they have been observed in nature at temperatures as cold as 2°C (Shapiro and Haugen, 1988), and 0°C (Gradinger and Lenz, 1995). Our deck incubation experiments showed that northern Synechococcus populations can actually grow at 2°C, although with a quite low growth rate (maximum of $0.13 d^{-1}$), suggesting a physiological adaptation of Arctic populations to low temperatures that further supports the existence of Synechococcus thermotypes (Pittera et al., 2014). This hypothesis is strengthened by our findings that many MicroPolar sequences formed new OTUs, unveiling an important novel genetic diversity (especially within clade I), which seems to be specific to this geographic area (17 OTUs out of the 41 OTU identified within clades I and IV). Furthermore, sequences obtained from August and November are mainly found in two OTUs within subclade Ib, gathering reference sequences retrieved only at high latitude from the Barents Sea (72°N) and the North Atlantic Ocean (57°N), but hardly detected in the Tara Oceans dataset. Altogether, these results point toward the existence of Synechococcus populations endemic to these Arctic or subarctic areas.

The peak-values of *Synechococcus* were clearly associated with the Atlantic inflow (salinity > 34.9) and abundances decreased exponentially with decreasing temperature and were

most often low in ice-associated water. This, along with the tendency of decreasing concentrations with decreasing salinity, is in accordance with the suggestion of Synechococcus being an indicator of saline Atlantic water transported into the Arctic (Murphy and Haugen, 1985; Gradinger and Lenz, 1995) as well as the low tolerance to wide salinity ranges of obligate marine Synechococcus (Waterbury et al., 1986). It should also be noted that although Synechococcus peak abundances were found in the relatively warm, saline Atlantic water, equally high abundances were observed in discrete samples from non-Atlantic water masses throughout the year (Figure 3), indicating the potential of Synechococcus to adapt to cold, low saline water, as also suggested by Nelson et al. (2014) for Canadian Arctic Synechococcus. The observed maximum abundance of picoeukaryotes, on the other hand, was found at a salinity of 33.5 and they were in general less affected by low salinities than Synechococcus. The dominance of picoeukaryotes over Synechococcus in the Arctic region may thus be connected to their capacity to stand a wide range of salinities in addition to an adaptation to low temperature. As only a few of our samples had a low salinity (17 surface samples in August have salinity <33), more efforts are needed to confirm this trend. In the Canada Basin of the Arctic Ocean proper Synechococcus abundance of 60 cell mL⁻¹ was found at salinities substantially lower than 33 (Li et al., 2013).

The extreme changes in light conditions in polar environments may also have been a driver for the diversification of the Synechococcus populations. However, in contrast to Prochlorococcus, obvious light partitioning is usually not observed for Synechococcus (Scanlan et al., 2009) since only one study reported a vertical partitioning of some Synechococcus genotypes so far (Gutiérrez-Rodríguez et al., 2014). In our incubations Synechococcus surprisingly showed a net growth in January and March when light was absent or low, respectively, while picoeukaryotes did not grow (data not shown) (Figure 6). The ability of Synechococcus to grow under very low light conditions is presumably related to their capacity to consume dissolved organic matter (Palenik et al., 2003; Cottrell and Kirchman, 2009). Yelton et al. (2016) indeed found that the genetic potential for mixotrophy in picocyanobacteria (through osmotrophy) is globally distributed. Although this still needs to be confirmed by laboratory experiments, it is possible that Synechococcus OTUs detected in November, when there is no light, belong to mixotrophic populations that are adapted to slow growth in the dark. Picoeukaryotes may use another mixotrophic strategy, i.e., bacterial grazing, to sustain growth during dark months (Sanders and Gast, 2012). Our observations that Synechococcus can be more abundant than picoeukaryotes in the Arctic in autumn and winter (Figure 4) are consistent with previous results [Gradinger and Lenz, 1995; unpublished results from Adventfjorden, Svalbard (I. Kessel Nordgård, personal communication)] and may suggest that cyanobacterial osmotrophy is a more efficient strategy than picoeukaryotic phagotrophy to survive in the dark.

Grazing on Synechococcus

The highest *Synechococcus* abundances were observed when NO_3^- concentrations were low. Hence, there is no reason to believe that they were resource controlled. The tendency of

increased growth when potential grazers were removed, rather points at a top-down control. The all-year-round presence of heterotrophic flagellates (HNF), considered to be their main predators (Sanders et al., 1992; Christaki et al., 2001; Kuipers et al., 2003; Zwirglmaier et al., 2009) indeed allows for grazer control of the Synechococcus populations. Still, grazing losses of Synechococcus are challenging to estimate as potential grazers can include various nano-but also microzooplankton and the specific loss also depends on the presence of other prey types (i.e., bacteria and picoeukaryotes; Pernthaler, 2005). This is illustrated by the different outcomes of successively removing various grazer fractions, which in March, August and November did not result in different growth patterns, but in January and May led to higher growth rates of Synechococcus when organisms larger than 90 μm were removed (Figure S1). Thus, this may reflect a trophic cascade where the microzooplankton graze on HNF and thereby release picoplankton from grazing pressure in the $<90 \ \mu m$ fraction. In March, August and November, however, there was little effect of size fractionation, which indicates that small HNF $(<3 \,\mu\text{m})$ were the main grazers of picoplankton and that these were not grazer-controlled themselves. Exactly "who" were the most important Synechococcus grazers is not possible to deduce from the presented data, and probably varies over the season. In addition, infection by viruses probably also functions as a top down regulator of these Synechococcus populations (Sandaa and Larsen, 2006), however virus counts remained relatively constant in all five experiments (data not shown). Still, we did find the highest net growth rates for Synechococcus when the HNF abundance was lowest (January and March) as well as the highest Synechococcus in situ abundance in water with low HNF concentration (and vice versa), which is in accordance with the view that HNF control their abundance and distribution at large. The picoeukaryote abundance did not follow the same patterns (data not shown), suggesting that they may have different predators. The fact that autotrophs, such as Synechococcus and picoeukaryotes, persist during winter in very low abundances further suggests that low encounter rates between predator and prey in the highly diluted wintry environment release the picophytoplankton from grazing pressure and allows survival despite adverse growth conditions (Kiørboe, 2008). The experiments also illustrate that Synechococcus in both January, March and August have the highest growth rates in the fractions where the total prey:HNF ratio is highest, indicating that Synechococcus might escape the grazers when other potential prey organisms are relatively abundant.

Synechococcus As an Active Player in the Arctic and Future Implications

It may be questioned whether the observed occurrence of *Synechococcus* was simply a result of advection and passive transport via the Atlantic water inflow. Since the highest measured abundances were found within the core of the Atlantic water, this probably represents the major source. The seasonal maximum *Synechococcus* abundance, which was observed in August, does however coincide in time with the seasonal *Synechococcus* bloom further south along the Norwegian coast. Given the average transportation time is at its minimum in summer (Fahrbach et al., 2001), it seems unlikely that the

encountered seasonal change in *Synechococcus* community we observed was a mere product of advection of Atlantic water. Moreover, the spatial and temporal distribution of clades and OTUs as well as the observed growth at low temperatures when released from grazing pressure, rather suggests that at least some of the observed *Synechococcus* populations are adapted to Arctic conditions and are indigenous to these waters.

Due to their small size $(1.1 \pm 0.4 \ \mu m$ diameter in the subarctic Atlantic; Paulsen et al., 2015), Synechococcus cells are largely grazed by HNF and microzooplankton (Christaki et al., 1999, 2005). This implies that their biomass production will be largely recycled in the microbial food web and thus be of minor contribution to higher trophic levels in the grazing food web. Even at the highest abundances observed in this study, Synechococcus only constitutes a minor part of the Arctic epipelagic carbon and energy pool (e.g., 21,000 cells mL^{-1} is equal to 2.3 μg C $L^{-1},$ assuming a diameter of 1.1 μm and 250 fg C μ m⁻³; Kana and Glibert, 1987) relative to the total phytoplankton biomass of 42 μ g C L⁻¹ (assuming a carbon to chl a conversion of 30). A warmer Arctic ocean that may favor Synechococcus at the expense of larger phytoplankton species (Flombaum et al., 2013) implies that more energy and carbon could be retained within the microbial food web, further reducing the contribution of Arctic primary production to the top of the food chain.

AUTHOR CONTRIBUTIONS

MP led the collection and analysis of data, and the writing of the paper. All other authors contributed to writing the paper and in addition AL, OM, RS, and LS helped collecting data and performing experiments. LG, HD, OM, and GB helped analyse the data and prepare figures.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmars. 2016.00191

Figure S1 | The abundance (cells mL⁻¹) of *Synechococcus* (red) and HNF (blue) plotted on the left y-axis during the first 5 days of fractionation experiments performed during the 5 cruises. The fractions <90 μ m (A-E), <10 μ m (F-J), <5 μ m (K-O) and <3 μ m (Q-U) are represented on each row. Exponential functions were fitted (lines) to the abundance providing the net growth rates (μ) given in the upper left corner for *Synechococcus* (red) and HNF (blue). The total prey (sum of *Synechococcus*, picoeukaryotes and heterotrophic bacteria) to HNF ratio is plotted for each triplicate on he right y-axis (open black circles), the black line connects the daily average prey:HNF ratio.

Figure S2 | The abundance (cells mL⁻¹) of *Synechococcus* (red) and picoeukaryotes (green) for all months within the upper 500 m, except for March where profiles are shown down to 1000 and 3000 m. Horizontal light blue lines mark the stations that were influenced by sea ice. Note the different x-axis for different months. Coordinates are given for each station above each graph.

Table S1 | Environmental from the cruises containing: dates (mm/dd/yy), latitude and longitude of stations (decimal degrees), depth (m), flow cytometer counts of *Synechococcus*, picoeukaryotes, nanophytoplankton, heterotrophic bacteria, and nanoflagellates (cells mL⁻¹), the growth rates *Synechococcus* and HNF (d⁻¹) from the <90 μ m incubation, salinity, temperature and potential temperature (°C), CTD-fluorescence (RUF), total chl a and the chl a fraction > 10 μ m (μ g L⁻¹), and nutrients (NH⁴₄, NO⁻₃, NO⁻₂, PO⁺₄, Si(OH)₄ (μ M). N.B. nutrients from January, May and August are not included here but will be available in Randelhoff et al. submitted.

 Table S2 | Sequence ID of the members of each Operational Taxonomical

 Unit (OTU) defined for *petB* at 97% nucleotide sequence identity.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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