INTRODUCTION

Sea ice provides a low-temperature habitat for diverse communities of microorganisms including viruses, bacteria and heterotrophic (e.g. flagellates and ciliates) and autotrophic protists (e.g. diatoms) (Kaartokallio et al. 2006). During sea ice formation, microorganisms, inorganic solutes and solids can be incorporated into the ice and accumulate to concentrations higher than that in the underlying seawater (Reimnitz et al. 1992, Grossmann & Gleitz 1993, Gradinger & Ikaävalko 1998). Organisms incorporated into sea ice are challenged with changes in space, light availability, salinity, nutrients, dissolved inorganic carbon (DIC) and O$_2$ concentration, temperature and pH (Gradinger & Ikaävalko 1998). Especially, light availability within the sea ice has a major influence on the sea ice algal biomass and production (Cota & Horne 1989, Rysgaard et al. 2001).

Sea ice algae constitute an important component of sympagic communities and have been extensively studied in Arctic sea ice (e.g. Horner & Schrader 1982, Gosselin et al. 1997, Glud et al. 2007, Mikkelsen et al. 2008). The sea ice algae represent an important food source for metazoan grazers, and photosynthetic prod-

ABSTRACT: We present a study of autotrophic and heterotrophic activities of Arctic sea ice (Malene Bight, SW Greenland) as measured by 2 different approaches: (1) standard incubation techniques (H$^{14}$CO$_3^-$ and [3H]thymidine incubation) on sea ice cores brought to the laboratory and (2) cores incubated in situ in plastic bags with subsequent melting and measurements of changes in total O$_2$ concentrations. The standard incubations showed that the annual succession followed a distinctive pattern, with a low, almost balancing heterotrophic and autotrophic activity during February and March. This period was followed by an algal bloom in late March and April, leading to a net autotrophic community. During February and March, the oxygen level in the bag incubations remained constant, validating the low balanced heterotrophic and autotrophic activity. As the autotrophic activity exceeded the heterotrophic activity in late March and April, it resulted in a significant net oxygen accumulation in the bag incubations. Integrated over the entire season, the sea ice of Malene Bight was net autotrophic with an annual net carbon fixation of 220 mg C m$^{-2}$, reflecting the net result of a sea ice-related gross primary production of 350 mg C m$^{-2}$ and concurrent bacterial carbon demand of 130 mg C m$^{-2}$. Converting the O$_2$ net exchange of the bag incubations into carbon turnover estimated an annual net carbon fixation of 1700 ± 760 mg C m$^{-2}$ (mean ± SD), which was higher than the annual net carbon fixation quantified in the standard incubations.

KEY WORDS: Sea ice · Primary production · Bacterial carbon demand · Net autotrophic activity · Net heterotrophic activity · Attenuation coefficients
ucts or entrained organic material can lead to elevated bacteria abundance and production within the sea ice (Smith et al. 1989, Kremsb et al. 2002, Meiners et al. 2003, Riedel et al. 2007).

Several previous studies have shown that heterotrophic bacteria are active and abundant in Arctic sea ice (Bunch & Harland 1990, Smith & Clement 1990, Brinkmeyer et al. 2003, Lizotte 2003, Kaartokallio 2004), but studies on spatial and seasonal variations are few (Smith et al. 1989, Gradinger and Zhang 1997).

Sea ice represents a partially interconnected network of brine-filled channels comprising 1 to 30% of the sea ice volume (e.g. Weeks & Ackley 1986). The degree of interconnection of the brine enclosures is generally enhanced with increasing temperature, and the potential accumulation of sea ice algae therefore increases towards the polar spring. The significance of heterotrophic processes typically increases during late bloom and postbloom situations close to spring thaw (Vézina et al. 1997, Kaartokallio 2004). In addition to biologically mediated dynamics, thaw and freezing processes induce extensive dynamics in solute and gas distribution (Glud et al. 2002). Consequently, the sea ice matrix is highly heterogeneous and dynamic, and quantification of in situ algae and bacterial productivity represent a true challenge.

The overall objective of this investigation was to describe the dynamics of autotrophic and heterotrophic activity in first-year sea ice. We measured autotrophic and heterotrophic activity of intact sea ice cores under in situ conditions in bag incubations, and measurements were compared with primary production and bacterial carbon demand as measured in the laboratory by standard H14CO3– and [3H]thymidine incubations under well-defined conditions.

**MATERIALS AND METHODS**

Measurements were conducted on first-year land-fast sea ice in Malene Bight in the vicinity of Nuuk, SW Greenland (64°82'N, 51°42'W). Sampling was performed at 1 to 2 wk intervals from 1 February to 14 April 2008. During that period, sea ice thickness varied between 0 and 62 cm, while snow cover ranged between 0 and 28 cm. The net aerobic activity of an enclosed sea ice community was followed in situ by determining the O2 concentrations in sea ice cores sealed in plastic bags and placed under natural snow cover, hereafter referred to as ‘bag incubations’. These measurements were supplemented with standard measurements of temperature, salinity, irradiance attenuation in snow and ice, nutrient concentrations (phosphate, PO43–; nitrate and nitrite, NO3– + NO2–; silicic acid, Si(OH)4; and ammonium, NH4+), chlorophyll a (chl a) concentrations, sea ice algal productivity and bacterial carbon demand during each sampling campaign, the latter 2 being referred to as ‘standard incubations’.

On 15 February and 19 March, 10 sea ice cores were collected using a MARK II coring system (Kovacs Enterprises). Cores were cut into 2 sections of equal length (ca. 22 cm), i.e. top half and bottom half, which were brought back to the laboratory in black plastic bags within 1 h of sampling. In a laboratory cold room (3 ± 1°C), core sections were placed in laminated transparent NEN/PE plastic bags (Hansen et al. 2000) fitted with a gas-tight Tygon tube and a valve for sampling.

Artificial seawater (salinity of 33) with a known O2 concentration was added (10 to 30 ml) to the NEN/PE bags (Hansen et al. 2000). The bags were closed, and excess air quickly extracted through the valve. The top and bottom halves of a single ice core (i.e. the contents of one pair of bags) were melted within 48 h in darkness (3 ± 1°C). Gas bubbles released from the melting sea ice were subsequently transferred to 12 ml Exetainer tubes (Labco) containing 20 μl HgCl2 (5% w/v, saturated solution). The gas bubbles were analysed for gaseous O2 by gas chromatography in a flame ionization detector and thermal conductivity detector (FID/TCD, SRI model 8610C). The melted water was also transferred to Exetainer tubes for O2 measurements. Dissolved O2 in the melted sea ice was measured by Winkler titration (Grasshoff et al. 1983). The remaining 9 sea ice cores in plastic bags were transferred to the drilled holes at the sea ice sampling site within 2 h, and the in situ snow cover above the cores was re-established. The sea ice cores, i.e. top and bottom halves, were sampled at 1 to 2 wk intervals to determine the O2 bulk concentration of the sea ice.

Permeability of the NEN/PE bags was quantified by adding artificial seawater (salinity of 33) to 9 plastic bags containing a gas-tight valve for sampling (Hansen et al. 2000). The bags were spiked with HgCl2 (200 μl of a 5% solution per liter seawater) to prevent biological activity during the incubation. The water was then flushed with N2 gas until the O2 concentration in the bags reached 50% of atmospheric saturation. The valves in the bags were then closed ensuring that no gas phase was left inside the 1 l bag. Three bags were incubated at –20°C, 3 bags at 3°C and 3 bags at 20°C for 7 d. Initially the O2 concentration was measured in all bags. After the incubation, O2 samples were collected and measured as described previously. Ice formation in the bags (–20°C experiment) led to bubble formation and the O2 concentration was therefore calculated as the sum of O2 in gas and water. The maximum O2 flux across the plastic bags was calculated from the O2 concentration change between initial and final readings to 0.48 ± 0.20 μmol O2 l–1 d–1 (mean ± SD). This put a lower limit on the activity that can be
resolved by this procedure and may compromise overall net activity at low biomass. In reality the permeability at the incubation temperature (–5 to 0°C) was significantly less than our maximum value.

O₂ bulk concentration in the sea ice (Cᵢ) was calculated as:

\[ Cᵢ = \frac{(Cₘ \times Wₘ) - (Cₐ \times Wₐ)}{Wₐ} \]  

(1)

where \( Cₘ \) is the O₂ concentration in the melted sea ice (gas bubble + melted sea ice), \( Wₘ \) is the weight of the melted sea ice, \( Cₐ \) is the O₂ concentration in the artificial sea water, \( Wₐ \) is the weight of the artificial sea water, and \( Wᵢ \) is the weight of the sea ice (Rysgaard & Glud 2004).

The season was divided into Series 1 and Series 2. Series 1 consisted of 10 cores, which were incubated on 15 February and were individually collected for analysis from 15 February to 25 March. Series 2 consisted of another 10 cores incubated on 19 March and were individually collected for analysis from 19 March to 11 April. Linear regression was performed on each of the 4 series (i.e. top half and bottom half of cores in Series 1 and Series 2). The slope of the regression line was tested in all 4 series by means of Student's t-test.

On each of the 7 sampling occasions, triplicate ice cores were collected using a MARK II coring system. The 3 sea ice cores were cut into sections using a stainless steel handsaw, and vertical temperature profiles were measured in drilled holes to the centre of each section using a thermometer (Testo thermometer). Downwelling irradiance was measured directly above and below the snow with a data logger (LI-1400, Li-Cor Biosciences). In addition, air temperature was measured 2 m above the snow, and snow and sea ice thicknesses were determined using a measuring stick. The sea ice sections were placed in plastic containers in a dark insulated transport box (Thermobox) and brought back to the laboratory. Light attenuation of the sea ice samples was measured with a LI-1400 data logger in a dark thermally regulated room using a fibre lamp with an optic tungsten–halogen bulb. The LI-1400 data logger was calibrated against a pure chl a standard (Turner Designs). The remaining filtered sea ice was frozen (–18°C) for later analysis of PO₄³⁻, NO₃⁻ + NO₂⁻, Si(OH)₄ and NH₄⁺. Concentrations of NO₃⁻ + NO₂⁻ were measured by vanadium chloride reduction (Braman & Hendrix 1989). Concentrations of Si(OH)₄ and PO₄³⁻ were determined by spectrophotometric analysis (Strickland & Parsons 1972, Grasshoff et al. 1983). Concentrations of NH₄⁺ were determined by a fluorometric method (Holmes et al. 1999). The lower detection limit for the nutrient measurements were 0.002 μmol l⁻¹ for PO₄³⁻, 0.5 μmol l⁻¹ for NO₃⁻ + NO₂⁻, 0.002 μmol l⁻¹ for Si(OH)₄ and 0.10 μmol l⁻¹ for NH₄⁺ (lower detection limit is calculated using the t-value of 2.99 corresponding to a 99% confidence interval with df = 7). Conductivity of the melted sea ice sections was measured using a conductivity cell (Thermo Orion 3-star with an Orion 013610MD conductivity cell) and converted to bulk salinity (Grasshoff et al. 1983). Sea ice brine salinity was calculated as a function of temperature (Cox & Weeks 1983) and the brine volume as a function of bulk salinity, density and temperature. Brine volume was calculated according to Leppäranta & Manninen (1988) for temperatures greater than –2°C and according to Cox & Weeks (1983) for temperatures less than –2°C.

Primary production was determined in melted (melted within 48 h in dark conditions at 3 ± 1°C) sea ice water at 3 light intensities (42, 21 and 9 μmol photons m⁻² s⁻¹) and corrected with one dark incubation using the H¹⁴CO₃⁻ incubation technique (Steemann-Nielsen 1952). The sea ice samples were melted and subsequently acclimatised at 20 μmol photons m⁻² s⁻¹ for a few hours (>2 h). Then the sea ice samples were poured into 120 ml gastight glass bottles and 4 μCi of H¹⁴CO₃⁻ were added to each bottle. The bottles were incubated on a plankton wheel at 1 rpm for 5 h at 3 ± 1°C at the different light intensities. Illumination was provided by cool white fluorescent lamps with a spectrum close to natural sunlight (Master TL-D 36W/840, Philips) and irradiance was measured using a LI-1400 data logger. Incubations were terminated by adding 200 μl of 5% ZnCl₂ and subsequently filtered onto 25 mm GF/F filters (Whatman). The filters were placed in scintillation vials containing 200 μl 1 M HCl to remove labelled, unfixed inorganic carbon, and then extracted in scintillation liquid (PerkinElmer Ultima Gold) for 22 h and counted using a liquid scintillation analyser (TricCarb 2800 TR, PerkinElmer). Dissolved inorganic carbon (DIC) concentrations in melted sea ice were measured on a CO₂ coulometer as described by Rysgaard & Glud (2004). After liquid scintillation counting, counts were converted to potential primary production (under photoinhibition) (PPᵢ, μg C l⁻¹ h⁻¹) as:

\[ PPᵢ = \frac{DPM_{\text{activity}} \times \text{DIC}_{\text{sea ice}} \times F_{\text{decl}} \times M_a}{DPM_{\text{added}} \times t_{\text{inc}}} \]  

(2)
where DPM_{activity} is the \(^{14}\text{C}\) assimilated carbon corrected for assimilated carbon in the dark (disintegrations per minute [dpm] on filter), DIC_{sea ice} is dissolved inorganic carbon in melted sea ice (\(-450\, \mu\text{mol}\ \text{l}^{-1}\)), F_{isc} = 1.05 is the discrimination factor of algae assimilation of \(^{13}\text{CO}_2\) and \(^{14}\text{CO}_2\), M_o is the molar mass of carbon (12.01 \text{g mol}^{-1}), DPM_{added} is the specific activity of the \(^{14}\text{C}\) labelled medium (dpm m\(^{-1}\)) in which cells were labelled, and \(t_{inc}\) is the incubation time (5 h).

The potential primary production (under no photoinhibition) (PP, \(\mu\text{g C}\ \text{l}^{-1}\ \text{h}^{-1}\)) measured in the laboratory at different sea ice depths was plotted against the 3 laboratory light intensities, 42, 21 and 9 \mu\text{mol} \text{photons m}^{-2} \text{s}^{-1}, and fitted to the following function described by Platt et al. (1980):

\[\text{PP} = P_m \left[1 - \exp\left(-\frac{\alpha E_{\text{PAR}}}{P_m}\right)\right] \tag{3}\]

where \(P_m\) (\(\mu\text{g C}\ \text{l}^{-1}\ \text{h}^{-1}\)) is the maximum photosynthetic rate at light saturation, \(\alpha\) (\(\mu\text{g C}\ \text{m}^2\ \text{s} \ \mu\text{mol} \text{photons}^{-1} \text{l}^{-1} \ \text{h}^{-1}\)) is the initial slope of the light curve and \(E_{\text{PAR}}\) (\(\mu\text{mol} \text{photons} \text{m}^{-2} \text{s}^{-1}\)) is the laboratory irradiance. The photoadaptation index, \(E_k\) (\(\mu\text{mol} \text{photons} \text{m}^{-2} \text{s}^{-1}\)) was calculated as \(P_o/\alpha\).

\textit{In situ} downwelling irradiance was measured at ground level with a pyrometer (Kipp & Zonen, model CM21, spectrum range of 305 to 2800 nm) once every 5 min, and hourly averages were provided by Asiaaq (Greenland Survey). Hourly downwelling irradiance was converted into hourly photosynthetically active radiation (PAR) (light spectrum, 300 to 700 nm) after intercalibration \((r^2 = 0.99, p < 0.001, n = 133)\) with a LI-1400 data logger (Li-Cor). The \textit{in situ} hourly PAR irradiance was calculated at different depths, depending on sea ice and snow thickness, using the attenuation coefficients measured during the sea ice season.

\textit{In situ} primary production was calculated for each hour at different sea ice depths using hourly \textit{in situ} PAR irradiance (see Eq. 3). Total daily (24 h) \textit{in situ} primary production was calculated as the sum of hourly \textit{in situ} primary production for each depth. The depth-integrated net primary production was calculated using trapezoidal integration.

Bacteria production in melted sea ice samples was determined by measuring the incorporation of \(^{3}\text{H}\)thymidine into DNA. Triplicate subsamples (volume, \(V_{\text{filt}} = 0.01\) l) were incubated in darkness at 3 ± 1°C for 42, 21 and 9 \mu\text{mol} \text{photons} m^{-2} s^{-1} with a LI-1400 data logger (Li-Cor). The \textit{in situ} PAR irradiance was calculated at different depths, depending on sea ice and snow thickness, using the attenuation coefficients measured during the sea ice season.

DPM_{activity} was added to all the subsamples (Fuhrman & Azam 1982). Triplicate subsamples (volume, \(V_{\text{filt}} = 0.01\) l) were incubated in darkness at 3 ± 1°C in scintillations vials until filtration. Subsamples were filtered through 25 mm mixed cellulose ester filters (pore size 0.2 \mu m, Advantec MSF) and the filters were placed in scintillation vials. Scintillation vials were rinsed with 5 ml of 5% cold TCA. Subsequently, filters were rinsed 7 times with 1 ml of cold 5% TCA and then extracted in scintillation liquid (Ultima Gold, Perkin-Elmer) for 22 h and counted using a liquid scintillation analyser (Tricarb 2800, PerkinElmer).

Bacteria production (BP, \(\mu\text{g C}\ \text{l}^{-1}\ \text{h}^{-1}\)) was calculated as:

\[\text{BP} = \frac{\text{DPM}_{\text{sample}} \times \text{N}_{\text{cells}} \times \text{N}_c \times M_c}{\text{SA} \times t_{inc} \times V_{\text{filt}}} \tag{4}\]

where DPM_{sample} is the average dpm for the live treatment subtracted from the average dpm for the TCA-killed controls, \(N_{\text{cells}}\) is the conversion factor (2.09 \times 10^{18} \text{cells mol}^{-1} \text{3H}, according to Smith & Clement 1990), \(t_{inc}\) is the incubation period \(t_{inc} = 6\) h, \(V_{\text{filt}}\) is the volume of the subsamples \(V_{\text{filt}} = 0.01\) l, \(M_c\) is the molar mass of carbon and SA is the specific activity of the thymidine solution \((2.24 \times 10^{16} \text{dpm mol}^{-1})\).

\[\text{N}_c = \frac{\text{Cell}_{\text{size}} \times C_{\text{factor}}}{1000000} \tag{5}\]

where Cell_{size} is the average bacteria cell size (2.473 \mu m^2) (Smith & Clement 1990), and C_{factor} is the factor used to convert cell volume to carbon \((0.12 \text{pg C} \mu\text{m}^{-3})\) according to Smith & Clement (1990).

Bacterial carbon demand (BCD, \(\mu\text{g C}\ \text{l}^{-1}\ \text{h}^{-1}\)) was calculated as:

\[\text{BCD} = \frac{\text{BP}}{BGE} \tag{6}\]

where BP is the bacteria production and BGE is the bacterial growth efficiency of 0.5 according to Rivkin & Legendre (2001). To extrapolate bacterial carbon demand to \textit{daily in situ} carbon demand we assumed that the respiration was light-independent (multiply by 24). The depth-integrated bacterial carbon demand was calculated using trapezoidal integration.

On 23 February, 10 sea ice cores were collected along a 10 m long section to investigate the spatial (horizontal and vertical) variability of the biotic conditions (chl a, primary production and bacterial carbon demand) and the abiotic conditions (sea ice temperature, bulk salinity and brine volume) in the sea ice. The 10 sea ice cores were collected close to the plastic bag incubations at 1 m intervals. The cores were cut into 2 sections, i.e. top and bottom halves, and brought back to the laboratory in a dark Thermobox for further analysis as described.

To extend the evaluation of spatial variability, a large-scale investigation of horizontal variability was conducted on 29 February. Fifty-two sea ice cores were collected along a 367 m long transect to investigate the
heterogeneity of chl a, sea ice temperature, brine salinity and brine volume in the bottom of the sea ice. The snow and sea ice thickness was measured as described previously. Four sea ice cores were sampled at 20 cm intervals at position 1 m. The first core was cut vertically into 2 pieces at every sampling position. Sea ice cores were collected at distances of 1, 3, 5, 7, 9, 20, 31, 42, 53, 64, 165, 266 and ca. 367 m. All the ice cores were brought back to the laboratory in a dark Thermo box for determination of brine salinity, volume and chl a concentration as described.

Spatial autocorrelation (Legendre & Legendre 1998) was used to analyse the horizontal distribution of chl a, sea ice temperature, bulk salinity, snow thickness and sea ice thickness. We assume that the variability along one line is the same as along a perpendicular line. The autocorrelation was estimated by Moran’s I coefficients (Moran 1950, Legendre & Legendre 1998). This coefficient was calculated for each of the following intervals along the transect (distance classes): 0 to 0.50 m, 0.50 to 1.5 m, 1.5 to 2.5 m, 2.5 to 5.5 m, 5.5 to 10.5 m, 10.5 to 20.5 m, 20.5 to 50.5 m, 50.5 to 100.5 m and >100.5 m. The autocorrelation coefficients estimated by Moran’s I coefficient were tested for significance according to the method described in Legendre & Legendre (1998). A 2-tailed test of significance was used. The null hypothesis of random spatial distribution was rejected at the specified level of significance when an individual autocorrelation coefficient exceeded a critical value (positive or negative). A significance level of p < 0.05 was used.

RESULTS

Abiotic parameters

Temperatures within the snow cover varied from −13 to 0°C and the sea ice temperature from −5 to 0°C (Fig. 1a). Minimum temperatures of the sea ice were...
observed in February, after which the temperature gradually increased to maximum values just before sea ice break-up in mid-April. The high light reflectance and scatter from the snow cover caused strong light attenuation throughout the sea ice season, with average attenuation coefficients of $K_{\text{snow}} = 23$ m$^{-1}$ and $K_{\text{ice}} = 8$ m$^{-1}$ (Fig. 1b). Early in the sea ice season, irradiance at the bottom of the sea ice was low (0.03 to 0.15 μmol photons m$^{-2}$ s$^{-1}$) (Fig. 1c). Light availability increased along with increasing day length and declining snow cover, reaching a maximum downwelling irradiance of 76 μmol photons m$^{-2}$ s$^{-1}$ in the uppermost sea ice section and 7 μmol photons m$^{-2}$ s$^{-1}$ in the bottom on April 4.

The bulk salinity decreased over time with a maximum salinity of 9 in early March and a minimum salinity of 1 from late March and onward (Fig. 1d). Bulk salinity varied vertically within sea ice cores during winter, with lower values encountered in the bottom sea ice from mid-February until late March. From 25 March to 4 April, bulk salinity was lowest in the uppermost part of the sea ice. The relative brine volume increased throughout the sea ice season, with a maximum in late March in the uppermost section of the sea ice (Fig. 1e). In late March melting from the top of the sea ice was initiated, which resulted in high relative brine volumes and low bulk salinities in the uppermost part of the sea ice. During this period air temperatures varied between 0 and 7°C.

### Nutrient parameters

Bulk $\text{PO}_4^{3-}$ concentration was initially 0.05 to 0.20 μmol l$^{-1}$ (Fig. 2a), and increased to a maximum of 0.70 μmol l$^{-1}$ in March but subsequently decreased to 0.05 μmol l$^{-1}$ in April. Bulk $\text{Si(OH)}_4$ concentration remained constant between 1.8 and 2.3 μmol l$^{-1}$ in the sea ice from February to late March with lowest values encountered at the bottom of the sea ice. In late March, $\text{Si(OH)}_4$ concentration decreased rapidly to below 0.4 μmol l$^{-1}$ (Fig. 2b). The initial bulk concentration of $\text{NO}_3^- + \text{NO}_2^-$ was 3.2 μmol l$^{-1}$ at the top and 2.0 μmol l$^{-1}$ at the bottom of the sea ice (Fig. 2c) and decreased throughout the season, reaching a minimum of 0.5 μmol l$^{-1}$ in April. The initial bulk concentration of $\text{NH}_4^+$ was 4.0 μmol l$^{-1}$ at the top and bottom of the sea ice. During the sea ice season the $\text{NH}_4^+$ concentration
increased, reaching values of 6 to 12 μmol l⁻¹ at the bottom of the sea ice and 5.0 to 6.0 μmol l⁻¹ at the top of the sea ice (Fig. 2d).

Bulk nutrient concentrations for each sampling date were plotted as a function of bulk salinity and compared with the expected dilution line (according to Clarke & Ackley 1984). If values were below the line, depletion of nutrients occurred in the sea ice. If values were above the dilution line, production or net deposition of the solute took place. Plots of salinity–PO₄³⁻, salinity–Si(OH)₄, salinity–NO₃⁻ + NO₂⁻ and salinity–NH₄⁺ in sea ice were generally all above the dilution line (Fig. 3), but most explicitly for NH₄⁺, which clearly accumulated within the sea ice (Fig. 3d).

**Biotic parameters**

Sea ice profiles of the algal biomass, expressed as chl a, showed that the highest bulk concentration in the lower 15 cm of the sea ice cores was 2.80 μg l⁻¹ on 25 March (Fig. 4a). Subsequently, the algal biomass in the lower 15 cm of the sea ice cores decreased rapidly reaching a value of 1.50 μg l⁻¹ in April.

Sea ice primary production integrated for the sea ice profile increased throughout winter from 0.09 mg C m⁻² d⁻¹ (15 February) to 12.60 mg C m⁻² d⁻¹ (4 April) (Fig. 4b). The highest volume-specific primary production (bulk) of 40 μg C l⁻¹ d⁻¹ was encountered in the middle part of the sea ice profile in April.

The highest sea ice bacterial carbon demand of 27 μg C l⁻¹ d⁻¹ was encountered in the central ice at the onset of the melting period (Fig. 4c). However, a single peak of 9.00 μg C l⁻¹ d⁻¹ was observed on 10 March in the upper 10 cm section of the sea ice.

Depth integration of the activity reflected low autotrophic and heterotrophic productivity during winter, followed by a slightly net heterotrophic period in late February and March. Finally, a net autotrophic period was observed from late March until the end of the study period (Fig. 5). Integrated over the entire measuring season (i.e. from 15 February to 14 April) the sea ice of Malene Bight was net autotrophic. An annual net carbon fixation of 220 mg C m⁻² was calculated by subtracting the net result of a sea ice-related gross primary production of 350 mg C m⁻² from the bacterial carbon demand of 130 mg C m⁻².

**Bag incubations**

Oxygen levels during February and March indicated a low net oxygen accumulation in the top sea ice cores of 0.50 ± 3.00 μmol O₂ l⁻¹ d⁻¹ (mean ± SD) and in the bottom sea ice cores of 1.30 ± 5.00 μmol O₂ l⁻¹ d⁻¹. However, none of these values were significantly different from
zero (p > 0.05) (Fig. 6). Autotrophic activity exceeded heterotrophic activity in late March and April, resulting in a significantly high net oxygen accumulation in the bottom sea ice cores of 6.30 ± 2.30 μmol O₂ l⁻¹ d⁻¹ (p < 0.01) whereas no significant oxygen accumulation (0.80 ± 3.50 μmol O₂ l⁻¹ d⁻¹) was observed in the top sea ice cores.

Assuming a photosynthetic quotient of 1.00 CO₂ evolved per O₂ consumed, a net annual carbon fixation of 1700 ± 760 g C m⁻² was calculated for the bag incubations.

Fig. 4. (a) Chl a concentrations (μg chl a l⁻¹) in bulk sea ice. (b) Primary production (μg C l⁻¹ melted sea ice d⁻¹). (c) Bacterial carbon demand (μg C l⁻¹ melted sea ice d⁻¹) calculated according to Rivkin & Legendre (2001). The black dots represent triplicate measurements.

Fig. 5. Primary production (PP, mg C m⁻² d⁻¹) and bacterial carbon demand (BCD, mg C m⁻² d⁻¹) in bulk sea ice.

Fig. 6. Measurements of O₂ concentration (μmol O₂ l⁻¹ melted sea ice), primary production (PP) (μg C l⁻¹ melted sea ice d⁻¹) and bacterial carbon demand (BCD) (μg C l⁻¹ melted sea ice d⁻¹) in (a) top half and (b) bottom half of sea ice cores. Dashed lines represent regression lines.

**Heterogeneity**

On 23 February the abiotic and biotic conditions, i.e. bulk salinity, brine volume, temperature, chl a, primary production and bacterial carbon demand, were measured in top and bottom sections of the sea ice cores (n = 10) along a 10 m transect (Table 1). There
was a significant effect of depth on chl $a$ levels (2-way ANOVA: $F_{1,19} = 7.452, p = 0.023$) with the bottom section having a higher phototrophic biomass than the top section of the ice cores. The primary production rates were also highest in the bottom sections and lowest in the top sections (2-way ANOVA: $F_{9,19} = 119.180, p < 0.0005$), while the bacterial carbon demand was higher in the top sections than in the bottom sections (2-way ANOVA: $F_{1,19} = 6.521, p < 0.031$) (Table 1).

Along a 10 m transect there was no significant horizontal variation across all biotic and abiotic conditions (2-way ANOVA: $F_{9,19} = 0.142, p \geq 0.142$). However, for all the parameters there was a significant vertical variability (2-way ANOVA: $F_{5, 49–50} = 15.215, p < 0.0005$).

Performing a similar statistical analysis on the temporal variations showed that, for all conditions, there was a significant temporal variability (2-way ANOVA: $F_{6,49–50} = 37.157, p < 0.0005$). Thus, for all conditions the horizontal variability in February was remarkably smaller than the seasonal variability.

Algal biomass (i.e. as chl $a$), bulk salinity and temperature in the lower section (5 cm) of the sea ice, and snow and sea ice thickness were measured along a 367 m transect to assess the horizontal distribution of these parameters. Moran’s $I$ was used to estimate the spatial autocorrelation within the data set (Moran 1950, Legendre & Legendre 1998, Rysgaard et al. 2001). Positive or negative values indicated positive or negative autocorrelation, respectively. The change from positive to negative values of Moran’s $I$ for chl $a$ occurred from the 20.5 m distance class to the 50.5 m distance class (Fig. 7). This indicates that the average sea ice algae biomass patch radius was between 20.5 and 50.5 m. The change from positive to negative values of Moran’s $I$ for sea ice temperatures and bulk salinity also occurred from the 20.5 m class of distance to the 50.5 m class of distance. In contrast, the average sea ice thickness, snow thickness and relative brine volume patch radius was >100.5 m, i.e. where the first change from positive to negative values of Moran’s $I$ occurred.

### DISCUSSION

Sea ice algae are known to be adapted to low ambient light levels and able to grow at light intensities as low as 7 to 20 μmol photons m$^{-2}$ s$^{-1}$ (Gosselin et al. 1990, Gradinger & Ikävalko 1998). Before April, snow cover inhibited any significant sea ice primary production (<1.5 mg C m$^{-2}$ d$^{-1}$) due to high attenuation coefficients and low light availability (<7 μmol photons m$^{-2}$ s$^{-1}$). As snow cover diminished, light availability increased to above 7 μmol photons m$^{-2}$ s$^{-1}$ and sea ice algae began to flourish; thus, the community became net autotrophic as light availability increased late in the season.

The annual succession of the sea ice organisms in Malene Bight followed a distinctive pattern, with a winter stage in February and March characterised by low and almost balanced heterotrophic and autotrophic activities. During February and March, the oxygen level in the bag incubations remained constant, validating the low, balanced heterotrophic and autotrophic activity (Fig. 6). As light availability increased in late March and April the now net autotrophic community led to a significant net oxygen accumulation of $6.30 \pm 2.30 \mu$mol O$_2$ d$^{-1}$ in the bottom sea ice cores whereas no significant oxygen accumulation ($0.80 \pm 3.50 \mu$mol O$_2$ d$^{-1}$) was observed in the top sea ice cores.
To correctly assess primary production and heterotrophic activity it is essential to have insight into in situ light availability. In the present study, high snow reflectance caused strong light attenuation for most of the sea ice season. The snow attenuation coefficient varied (from 4 to 32 m\(^{-1}\)) throughout the sea ice season with the highest attenuation coefficient of 32 m\(^{-1}\) found in March (Fig. 1b). The snow attenuation coefficients in Malene Bight were within the range of <4 to 40 m\(^{-1}\) reported by Weller & Schwerdtfeger (1967) and Thomas (1963) for natural snow cover in the Arctic and Antarctic. Dry and fresh snow as observed in Malene Bight has a significantly higher light attenuation than did compressed or wet snow at subzero temperatures (Perovich 1996, Perovich et al. 1998, Glud et al. 2007), resulting in low in situ primary productivity as was observed in February and March.

The average attenuation coefficient of 8 m\(^{-1}\) in the sea ice of Malene Bight was also relatively high compared with the attenuation coefficient during the summer thaw as reported by Glud et al. (2007) and Lüthje et al. (2006). The air humidity above the ice in Malene Bight was, on average, only 70% and in areas with low air humidity, air pockets could form due to brine drainage from surface layer, resulting in a high degree of scatter and increasing attenuation coefficients (Perovich 1996, Andreas & Ackley 1982, Trodahl & Buckley 1990). Overall our data on snow and sea ice attenuations are generally higher than values reported elsewhere, and as snow cover regulates light availability and therefore spatial patchiness of ice algae (Gosselin et al. 1986, Rysgaard et al. 2001), the phototrophic rates obtained during the present study were low. Sea ice algal biomass (chl a) patch size followed the snow and sea ice temperature patch size with an average radius of 20.5 to 50.5 m (Fig. 7). This suggests that snow cover and temperature (via brine volume) in sea ice were the main factors controlling sea ice algal patchiness. However, snow cover, temperature and light availability are crosscorrelated, and all exert a regulating effect on sea ice algal biomass in Malene Bight.

The maximum depth-integrated sea ice algal biomass of 0.07 to 0.5 mg chl a m\(^{-2}\) (Fig. 4b) was >2 orders of magnitude lower compared with the values of 50 to 150 mg chl a m\(^{-2}\) that were reported by Arrigo (2003) to be common in Arctic sea ice. Likewise, primary production measurements were low during February and March but gradually increased during the season, reaching a peak activity of only 12.6 mg C m\(^{-2}\) d\(^{-1}\) in early April, which is low compared with other studies (Arrigo 2003 and references therein). Integrated over the entire measuring period the annual sea ice algal carbon production was 350 mg C m\(^{-2}\), which was 47% lower than measurements made in the neighbouring Kobbefjord during 2006 (Mikkelsen et al. 2008). Furthermore, the sympagic primary production of 350 mg C m\(^{-2}\) corresponded to <1% of the annual pelagic primary production of the region (Juul-Pedersen et al. 2009).

The low biomass and primary production values of the sea ice in Malene Bight can be attributed to relatively low light availability and a relatively thick dense snow cover, which was also observed in NE Greenland by Glud et al. (2007). However, nutrient supply and different physical conditions, e.g. ice temperature, brine salinity and flushing of sea ice brine, and biological interactions also regulate the distribution and activity of sea ice organisms. The sea ice contained relatively high concentrations of NH\(_4\)^+, while PO\(_4\)^3−, Si(OH)\(_4\) and NO\(_2\)\(^-\) + NO\(_3\)\(^-\) concentrations in the sea ice were low (0.70 to 0.05 μmol l\(^{-1}\), 2.30 to 0.02 μmol l\(^{-1}\) and 3.2 to 0.5 μmol l\(^{-1}\), respectively (Figs. 2 & 3). The algal biomass was low at the bottom of the sea ice after mid-March, suggesting that the sea ice biomass was nutrient-limited late in the sea ice season. Nutrient concentration in sea ice is affected by biological activity as well as by exchange of nutrients between the sea ice, water and atmosphere. Previous studies have shown that during sea ice formation, dissolved constituents (including inorganic nutrients) are rejected from the ice matrix (Clarke & Ackley 1984, Giannelli et al. 2001). The nutrient–salt plots in Fig. 3 indicate that some import or accumulation of nutrients occurred during February and March. However, late in the sea ice season nutrients were generally depleted (Fig. 3), presumably due to microbial uptake (Fig. 4), which correlated with high primary productivity and a net autotrophic sea ice.

Plots of salinity–NH\(_4\)^+ suggested NH\(_4\)^+ enrichment, which occurred during the entire season, indicating an input from the atmosphere and/or heterotrophic activity by bacteria mineralising organic material encapsulated during sea ice formation. The standard laboratory-based incubations indicate a low net heterotrophic activity in February and March (Figs. 4 & 5). Calculated average bacterial carbon demand in this period was 1.60 μg C l\(^{-1}\) d\(^{-1}\). At a C:N ratio of 7:1, this is equivalent to a remineralisation rate of 0.02 μmol N l\(^{-1}\) d\(^{-1}\), which is lower than the values of Riedel et al. (2007) who reported NH\(_4\)^+ regeneration rates of 0.1 to 1.2 μmol N l\(^{-1}\) d\(^{-1}\) corresponding to only 20% of the winter–spring increase in NH\(_4\)^+ concentrations in the sea ice of Malene Bight. This suggests that NH\(_4\)^+ regeneration rates were not the main source of NH\(_4\)^+ accumulation in the sea ice. Thus, additional atmospheric input could potentially cause the high NH\(_4\)^+ concentrations in the sea ice of Malene Bight. In previous studies, high NH\(_4\)^+ values in sea ice have been measured in Arctic and Antarctic sea ice caused by...
regeneration of nitrogen compounds within the sea ice (Thomas et al. 1995, Kaartokallio 2001, Riedel et al. 2007).

Bacteria production rates of 0.05 to 1.3 mg C m⁻³ d⁻¹ in Malene Bight compare with rates of 0.004 to 6.0 mg C m⁻³ d⁻¹ reported by Kottmeier et al. (1987), 0.2 to 10 mg C m⁻³ d⁻¹ by Grossmann & Dieckmann (1994) and 2.9 to 5.6 mg C m⁻³ d⁻¹ by Mock et al. (1997).

The annual net bacterial carbon demand of 130 mg C m⁻² was 37% of the co-occurring primary production of 350 mg C m⁻². This observation is similar to findings reported in the world’s oceans (Ducklow 1983) and in benthic systems (Fenchel & Glud 2000). However, the ratio reported in the present study is significantly higher than the few percent reported from Antarctic (Kottmeier et al. 1987) and Arctic sea ice (Smith et al. 1989, Smith & Clement 1990). Our bag incubations from winter, however, support results from our traditional standard incubations that a close connection exists between phototrophic and heterotrophic communities and, considering that both algae and bacteria are situated close together in the sea ice brine matrix, a tight metabolic coupling would be expected. Positive correlation has also been demonstrated in some sea ice habitats (e.g. Gradinger & Zhang 1997, Meiners et al. 2003). However, limited data on the coupling between phototrophic and heterotrophic communities in sea ice exists and further investigations are required to elucidate the metabolic coupling in sea ice.

In the standard incubations, primary production of sea ice algae was determined by the ¹⁴C method (Steevan-Nielsen 1952). One problem with this technique is that melting or crushing of sea ice is required to homogeneously distribute ¹⁴C in the media, which separates organisms from the surrounding sea ice and alters the complex sea ice microenvironment. The photosynthetic performance may be considerably affected when the sea ice environment (i.e. irradiance, substrate, salinity, temperatures and spectral composition) is altered. Secondly, melting of the sea ice may influence algal abundance and species diversity (Gradinger et al. 1999). In the present study the samples were melted slowly (48 h) in darkness at 3 ± 1°C. In general decreasing or increasing salinity from that typical of seawater results in decreasing photosynthetic rates. Kirst & Wiencke (1995) showed that most sea ice algae are more tolerant to reduced rather than elevated salinities. Salinity-induced stress to sea ice algae is light dependent, such that incubated samples only suffered photosynthetic damage when irradiance was applied (Ralph et al. 2007). Furthermore, a potential problem when melting sea ice samples in darkness is inactivation of photrophs, which might remain inactive during a subsequent ¹⁴C incubation period in light. However, sea ice algae can adapt to increasing light intensity within hours (Smith & Sakshaug 1990). In addition, Peters & Thomas (1996b) showed that some algae cells are able to sustain an active photosynthetic apparatus for months in darkness, which allows carbon assimilation immediately after reintroduction to light. Also, photosynthetic rates in sea ice algae can be influenced by temperature (Palmisano et al. 1987, Ralph et al. 2005). Optimum temperature for ice algal photosynthesis is between 2 and 15°C, with a rapid decline in photosynthetic rate at temperatures above 15°C and below −10°C (Bunt 1964, Palmisano et al. 1987, Kottmeier & Sullivan 1988, Ralph et al. 2005). In the present study the sea ice was slowly melted at 3 ± 1°C; thus, the primary production was measured at higher temperatures than in situ. This suggests that the primary production might be overestimated in the winter period, where the difference between in situ temperature and laboratory temperature was greatest. However, it is a nontrivial task to incubate samples at low bulk salinity at 0°C or subzero temperature without introducing freezing and thawing artefacts. Overall, it is difficult to assess the extent to which our (and other) procedures for ¹⁴C incubations result in under- or over-estimation of the in situ primary production.

While calculation of phototrophic activity is relatively straightforward, the assessment of the bacteria production and bacterial carbon demand is based on a number of debatable assumptions. The method assumes that all bacteria assimilate exogenous thymidine and that eukaryotes do not. Uptake of [³H]thymidine by eukaryotic microalgae has been clearly demonstrated (Rivkin 1986). However, Fuhrman & Azam (1980) have shown that at low concentrations and relatively short incubation periods the uptake of [³H]thymidine by eukaryotes should be negligible. The [³H]thymidine method is a widely used index of bacterial production but requires a conversion factor to relate DNA production to production of bacterial cells. In the present study, we extrapolated incorporation data to bacteria production by applying the early season data of Smith & Clement (1990). They calculated the conversion factor for high Arctic sea ice bacteria as 2.09 × 10¹⁸ cells mol⁻¹ in the early sea ice season and as 0.47 × 10¹⁸ cells mol⁻¹ in the late sea ice season. The conversion factor from the early sea ice season fell within the commonly reported range of 1 to 4 × 10¹⁸ cells mol⁻¹ (e.g. Fuhrman & Azam 1982, Riemann et al. 1987), and Smith & Clement (1990) suggested that this factor was more appropriate than the lower conversion factor from the late season. Other factors used for the conversion of thymidine incorporation into bacteria production include the average bacterial cell size (Cellsize) and the volume-specific carbon content (Cfactor), which combined provide the cell-specific carbon content (NC) and which are affected by environmental changes.
(Coveney & Wetzel 1988). Integrated over the entire measuring season, the sea ice of Malene Bight was net autotrophic. However, when using different N factors discussed in the literature (Riemann et al. 1987, Smith & Clement 1990, Gradinger and Zhang 1997, Kaar-tokallio et al. 2005), the annual bacterial carbon demand ranged by a factor of 6 from 70 to 400 mg C m⁻² depending on which Nₐ value is used. In the present study, we used an Nₐ value measured in the high Arctic (Smith & Clement 1990) and, thus, we observed an annual carbon demand of 130 mg C m⁻².

Rivkin & Legendre (2001) reported that the bacterial growth efficiency was an inverse function of temperature. We used a growth efficiency measured in polar oceans of 0.50 (Rivkin & Legendre 2001). However, small changes in temperature would influence growth efficiency and, hence, bacterial carbon demand (~2.5% decrease in growth efficiency per 1°C increase). Using the 3 different growth efficiencies of 0.10, 0.25 and 0.90 (Bjørnsen 1986, Bjørnsen & Kuparinen 1991, Gradinger & Zhang 1997, Middelboe & Søndergaard 1993, Rivkin & Legendre 2001 and references therein, M. Middelboe & R. Glud unpubl. data), the annual bacterial carbon demand ranged by a factor of 5 from 71 to 635 mg C m⁻² depending on the growth efficiency we used. The conclusion would change from an annual net autotrophic sea ice to a net heterotrophic sea ice if we used the growth efficiency of 0.10. However, when using the growth efficiencies of 0.25 and 0.90 the sea ice is still net autotrophic. According to Rivkin & Legendre (2001), growth efficiency increases with decreasing temperatures. Thus, the conversion of bacteria production to bacterial carbon demand by assuming a growth efficiency of 0.5 appears to be a realistic compromise among the conflicting literature values of 0.10 to 0.90 (Bjørnsen 1986, Bjørnsen & Kuparinen 1991, Gradinger & Zhang 1997, Middelboe & Søndergaard 1993, Rivkin & Legendre 2001 and references therein). It would be ideal to determine the respective factors at different seasonal conditions to improve the assessment of bacterial production and carbon demand, but in practice this would be an unrealistically large effort for most studies.

The O₂ concentration of sea ice is regulated by photosynthesis and respiration (Gleitz et al. 1995, Günther et al. 1999) and is also affected by freezing and thawing of sea ice (Glud et al. 2002). The temporal and spatial variations in snow cover thickness can introduce variability in sea ice temperatures and consequently in O₂ concentrations. During the present study, the snow cover varied from 0 to 26 cm and the sea ice temperature varied from 0 to ~6°C (Fig. 1a), and only slight changes in temperature can lead to leakage of supersaturated and subsaturated water from the sea ice matrix (Glud et al. 2002). It is generally accepted that to obtain realistic estimates of in situ microbial activity in sea ice, the microenvironment (i.e. temperature, salinity, nutrient concentration) should be maintained (e.g. McMinn & Ashworth 1998). The bag incubations were incubated in situ and, thus, the sea ice microenviornment was better maintained compared with standard incubations. This supports the use of bag incubations for determination of net O₂ production or consumption under natural conditions. In the present study, oxygen levels remained almost constant during February and March, indicating no or low net oxygen accumulation in the top sea ice cores of 0.50 ± 3.00 μmol O₂ l⁻¹ d⁻¹ and in the bottom sea ice cores of 1.30 ± 5.00 μmol O₂ l⁻¹ d⁻¹ (Fig. 6). The permeability of the bags was tested and could be responsible for a maximum O₂ exchange of 0.48 ± 0.20 μmol O₂ l⁻¹ d⁻¹, which may compromise the results in February and March when no or low net oxygen accumulation was observed.

The autotrophic activity exceeded the heterotrophic activity in late March and April, resulting in a high net oxygen accumulation in the bottom sea ice cores of 6.30 ± 2.30 μmol O₂ l⁻¹ d⁻¹, whereas no significant oxygen accumulation occurred in the top sea ice cores (0.80 ± 3.50 μmol O₂ l⁻¹ d⁻¹). The O₂ measurements showed high variability, indicating small-scale heterogeneity in oxygen. Thus, we suggest that more replicates be applied from all time intervals to quantify net oxygen consumption or production in future studies.

The temperature of sea ice enclosed in the bags did not differ from the ambient temperature profiles (data not shown). However, enclosure might have a number of consequences for the measured microbial activity. In case of high sea ice algal primary production and low nutrient concentrations, enclosing sea ice cores in bags creates a risk of underestimating primary production, as nutrient consumption sets an upper limit to production. In the present study, primary production measured in the bags from the bottom of the sea ice during peak activity late in the season was 60 μg C l⁻¹ d⁻¹. Assuming a C:N Redfield ratio of 6.6 this represents a sufficient nitrogen pool for only 7 d of incubation. This indicates that the enclosed primary producers could have been nutrient-limited during the late part of the season (Fig. 6b), which suggests that oxygen production in the bottom sea ice section in the final incubation was underestimated. However, the annual net carbon fixation in the bag incubations was 8-fold higher than the annual net carbon fixation of 220 mg C m⁻² quantified from the standard incubations. We cannot, on this basis, pinpoint the reason for the apparent mismatch or evaluate which estimate is most correct. However, bags incubated for <7 d create a minimum disturbance of the microenvironment within the sea ice and might be a strong complementary tool to standard techniques.
when assessing the net activity of microbial communities in sea ice. Hence, further work with the bag incubation technique in situ is clearly needed to better elucidate whether sea ice, on an annual scale, is net autotrophic or net heterotrophic.

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