Holocene variability in sea ice and primary productivity in the northeastern Baffin Bay

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Abstract

Arctic sea ice is a critical component of the climate system, known to influence ocean circulation, earth’s albedo, and ocean–atmosphere heat and gas exchange. Current developments in the use of IP25 (a sea ice proxy with 25 carbon atoms only synthesized by Arctic sea ice diatoms) have proven it to be a suitable proxy for paleo-sea ice reconstructions over hundreds of thousands to even millions of years. In the NE Baffin Bay, off NW Greenland, Melville Bugt is a climate-sensitive region characterized by strong seasonal sea ice variability and strong melt-water discharge from the Greenland Ice Sheet (GIS). Here, we present a centennial-scale resolution Holocene sea ice record, based on IP25 and open-water phytoplankton biomarkers (brassicasterol, dinosterol and HBI III) using core GeoB19927-3 (73° 35.26′ N, 58° 05.66′ W). Seasonal to ice-edge conditions near the core site are documented for most of the Holocene period with some significant variability. In the lower-most part, a cold interval characterized by extensive sea ice cover and very low local productivity is succeeded by an interval (~ 9.4–8.5 ka BP) with reduced sea ice cover, enhanced GIS spring melting, and strong influence of the West Greenland Current (WGC). From ~ 8.5 until ~ 7.8 ka BP, a cooling event is recorded by ice algae and phytoplankton biomarkers. They indicate an extended sea ice cover, possibly related to the opening of Nares Strait, which may have led to an increased influx of Polar Water into NE-Baffin Bay. The interval between ~ 7.8 and ~ 3.0 ka BP is characterized by generally reduced sea ice cover with millennial-scale variability of the (late winter/early spring) ice-edge limit, increased open-water conditions (polynya type), and a dominant WGC carrying warm waters at least as far as the Melville Bugt area. During the last ~ 3.0 ka BP, our biomarker records do not reflect the late Holocene ‘Neoglacial cooling’ observed elsewhere in the Northern Hemisphere, possibly due to the persistent influence of the WGC and interactions with the adjacent fjords. Peaks in HBI III at about ~ 2.1 and ~ 1.3 ka BP, interpreted as persistent ice-edge situations, might correlate with the Roman Warm Period (RWP) and Medieval Climate Anomaly (MCA), respectively, in-phase with the North Atlantic Oscillation (NAO) mode. When integrated with marine and terrestrial records from other circum-Baffin Bay areas (Disko Bay, the Canadian Arctic, the Labrador Sea), the Melville Bugt biomarker records point to close ties with high Arctic and Northern Hemispheric climate conditions, driven by solar and oceanic circulation forcings.

Keywords Holocene · Baffin bay · Sea ice · IP25 · HBI III · Brassicasterol · PIP25 index

Introduction

Polar regions in the Northern Hemisphere, especially the Arctic and Greenland areas, are undergoing dramatic changes due to an accelerating reduction in sea ice cover and ice sheet extent for the past three–four decades. Since the late 70s, the Arctic sea ice cover has decreased rapidly [57, 94]. The summer sea ice extent and thickness of multi-year ice are decreasing at the fastest rate ever observed during recent times (~ 8.6% per decade), and model simulations suggest that Arctic summer sea ice may
disappear within the next 50 or even 30 years [43, 114, 118, 130]. Given the magnitude of the ongoing reduction in the sea ice cover, information on its impact on water mass circulation and related natural climate conditions over longer time scales is important, notably for the testing of predictive climate models. However, historical and satellite records only span a few centuries or decades, respectively [94, 128], thus limiting the understanding of natural variability of sea ice, emphasizing the need for high-resolution multi-proxy records of its past variations [53, 81]. Reliable long-term, at least semi-quantitative sea ice reconstructions are, thus, needed for the validation of Arctic climate scenarios [25, 28]. Furthermore, the ongoing enhanced melting of the Greenland Ice Sheet (GIS), and its significant contribution (~ 0.34 mm year\(^{-1}\)) [102] to global annual sea-level rise (~ 3.3 mm year\(^{-1}\)) [23], also needs to be appraised in relation to natural processes that occurred at millennial to centennial time scales during the present interglacial.

Northern Baffin Bay (Fig. 1A) is of particular interest due to the interaction of the relatively warm high-salinity West Greenland Current (WGC) water with the cold polar water of the Baffin Current (BC) originating from the Arctic Ocean, resulting in the presence of the high productivity North Water Polynya (NWP) kept open by winds, tides and ice bridges [5]. Eastward of the polynya, Melville Bugt borders the heavily glaciated NW Greenland region, only separated by narrow fringes of ice-free islands, nunataks, and peninsulas. Fast-flowing (> 1000 m a\(^{-1}\)) glaciers such as Ussing Brae and Cornell glaciers are in close proximity to Melville Bugt [54, 102, 133]. Marine sedimentary archives from this area may, thus, provide records of sea ice and past ice sheet dynamics in this highly climate-sensitive area. So far, the sea ice variability and long-term changes in water masses of the NE-Baffin Bay, especially the WGC, as well as the impact of melt-water discharge from the GIS that occurred since deglaciation, all remain barely documented [18, 22, 41].

The present study aims at filling this gap. It is based on a detailed analysis of a ~11.5-m-long gravity core from the southern Melville Bugt (core GeoB19927-3; Fig. 1A) spanning a time interval of the last approximately 10 ka BP. Special attention is paid to the high-resolution reconstruction of sea ice conditions, mainly documented from the abundance and flux of IP25 and open-water phytoplankton biomarkers (brassicasterol, dinosterol and HBI III) and sand content (grain size). These data are complemented by information about the core lithology and dry bulk density, as well as additional published paleoclimate records from circum-Baffin Bay areas.

**Biomarker proxies for paleo-environmental reconstruction**

Paleo-sea ice reconstructions rely heavily on proxy-based methods. In this context, combination of proxies (i.e., sedimentological, biogeochemical and micropaleontological) preserved in marine sediments such as ice-rafted debris, assemblages of diatoms, dinocysts and ostracods as well as \(\delta^{18}O\) in foraminiferal tests may provide indirect information about past sea ice conditions (e.g., [27–29, 40, 42, 50, 73, 83, 98, 131]). However, microfossil records such as, for example, diatom assemblages must be interpreted with caution as they may also be influenced by dissolution and selective preservation [14, 72, 110].

As a more direct approach, the “sea ice proxy IP25”, after its first introduction in a pioneer study by Belt et al. [7], has been preferentially used for reconstructing the variability of sea ice in the Arctic region (see [10] for a recent review). IP25 is a highly branched isoprenoid (HBI) alkene with 25 carbon atoms, derived from specific sea ice diatoms (mostly *Haslea* spp.) living in/under first-year ice and in brine channels [7, 8, 20]. As IP25 preserves well in the sediments and is resistant to degradation, it was possible to reconstruct sea ice variability dating back to hundreds of thousands to even millions of years ago [58, 65, 116, 117].

Further, an even more reliable and detailed picture is achieved when combining IP25 with open-water phytoplankton biomarkers such as brassicasterol, dinosterol and/or a tri-unsaturated HBI (Z-isomer; hereinafter referred to as “HBI III”) and its E-isomer [9, 12, 87, 89, 90, 112, 134]. In this way, the problem of distinguishing perennial ice cover from ice-free water can be solved. The absence or low values of both IP25 and phytoplankton biomarkers indicate a more permanent ice cover, whereas a minimal (to zero) IP25 and high phytoplankton biomarkers are indicative of ice-free
conditions, and a variable amount of these biomarkers may reflect changing seasonal sea ice cover. This approach has been successfully applied to sediments from the Fram Strait, the central Arctic Ocean, and Arctic marginal seas to reconstruct sea ice conditions during late Quaternary times (e.g., [12, 13, 36, 87, 88]), for the most recent review with pros and cons of this approach see [10].

The use of different phytoplankton biomarkers (in combination with IP25) may even allow distinguishing between different seasonal sea ice conditions including marginal ice zone (MIZ) situations (e.g., [9, 12, 103]). Based on the distribution pattern of HBI III in surface sediments from the western Barents Sea and the correlation with seasonal sea ice concentration maps, for example, Belt et al. [12] proposed that HBI III maxima reflect winter MIZ conditions. Based on a study of East Greenland fiords surface sediments, on the other hand, Ribeiro et al. [103] proposed that HBI III maxima correlate with the July MIZ situation. Based on biomarker investigations of surface sediments across Baffin Bay and the correlation with modern satellite observations, Kolling [61] suggested that the sea ice indices P2IP25 and P8IP25 may record the late spring and/or autumn conditions, whereas P8IP25 index may reflect more the late winter/early spring (ice-edge) conditions in the Baffin Bay. Furthermore, Belt et al. [9] recently suggested that the proportions of tri-unsaturated HBIs; Z and E-isomer (TR25) may indicate spring phytoplankton blooms, at least for the Barents Sea region. This approach, however, needs to be further evaluated using additional data from surface sediments and sediment cores from other Arctic regions.

**Environmental setting**

Melville Bugt is part of the broad shelf area in NE Baffin Bay (Fig. 1A) covering ca. 120,000 km² in the 200-km-wide continental shelf offshore NW Greenland. Baffin Bay acts as a pathway for sea ice and fresh-water exchange between the Arctic and North Atlantic Oceans. The oceanographic circulation in Baffin Bay (Fig. 1A) is characterized by the WGC (~150–1330 m water depth), flowing northwards along the west coast of Greenland and the cold BC (~100–300 m water depth) flowing southward and originating from the Arctic Ocean [119]. The WGC is formed by a combination of: (1) Atlantic sourced, relatively warm and saline water from the Irminger Current (IC), (2) polar sourced cold, low salinity water from the East Greenland Current (EGC) and (3) local melt-water discharge along the SW Greenland coast (Fig. 1A) [119]. The WGC near Melville Bugt is believed to split into two branches, one turning west that joins the BC near Smith Sound and the other branch circulating towards Nares Strait [21, 105].

This area displays high seasonal sea ice variability, especially the Melville Bugt area off West Greenland (Fig. 1B). In winter, this area is almost completely ice-covered as the sea ice margin, the so-called ‘Westice’, expands further southwards [113, 119]. In summer, however, the area becomes almost ice free. The ecological setting of Melville Bugt is highly affected by outlet glaciers of the GIS and receives about ~27% of GIS drainage [102]. Wind stresses here are highly seasonal with stronger wind stress in winter compared to summer [119].

During the Last Glacial Maximum (LGM), the extended Greenland, Laurentide, and Innuitian ice sheets (GIS, LIS, respectively) stretched across Baffin Bay coast to the shelf edge as demonstrated by the presence of cross-shelf troughs [38, 92, 111]. Although no geochronological records are currently available for the onset of deglaciation in Melville Bugt, numerical modeling data suggest the onset of deglaciation occurred as early as at ~16.0 ka BP [69]. However, the general-present day like ocean mass circulations were probably established in the central-eastern Baffin Bay ~14 ka BP and by ~10.4–9 ka BP in northern Baffin Bay [71, 109]. The opening of Nares Strait ~9 ka BP allowed the connection between Baffin Bay and the Arctic Ocean, which led to the establishment of the modern ocean circulations in the Baffin Bay [39, 48, 52]. The sediments deposited at the seafloor of Baffin Bay are predominantly derived from glacial erosion of the surrounding land masses, and high sedimentation rates of 40–140 cm ka⁻¹ make this area an ideal site for paleoenvironmental studies [115].

**Material and methods**

**Field methods**

The current study is based on the analysis of the 1147-cm-long gravity core GeoB19927-3 (Fig. 1A; black circle) (Lat: 73° 35.26′ N; Lon: 58° 05.66′ W; Water depth: 932 m), recovered from southern Melville Bugt during the RV Maria S. Merian cruise MSM44 (BAFFEAST) in June/July 2015 [31]. The working half of the core was sampled and divided into four sets of samples. One set was freeze-dried for biomarker analysis (5–10-cm resolution in this study) and the second set was used for dating (foraminifera and 210Pb). The other two sets were stored at +4 °C for other multi-proxy analyses (i.e., dinoflagellates, foraminifera, and provenance studies).

**Chronology/age model**

Age control is provided through a combination of radiocarbon and 210Pb-dating, followed by Bayesian age modeling.
Accelerator Mass Spectrometry (AMS) 14C-dating was performed on 12 samples, partially on mollusc shell fragments, partially on mixed benthic foraminifera (in one case mixed with specimens of the planktic foraminifera *Neogloboquadrina pachyderma* sin.) (see Table 1 for details). Mollusc fragments were measured at the Poznan Radiocarbon Laboratory, Poland; whereas, the foraminifera were dated at the MICADAS facility at the Alfred-Wegener-Institute in Bremerhaven, Germany. The larger mollusc fragments were measured as graphite, whereas a novel technology for very small sample sizes was used for the foraminifera picked from the > 100 μm fraction. Here, the CO2 released from the foraminiferal carbonate upon acid treatment is directly analyzed with a compact AMS facility equipped with a hybrid ion source (for a detailed explanation, see [127]). Radio-nuclide analyses (210Pb, 40K, 137Cs) were performed at the Bremen State Radioactivity Measurements Laboratory on a total of seven one-cm slices of sediment.

The final age model was constructed using the software package *BACON* [15], written as open-source code for the statistical computation program “R”. BACON uses a Bayesian approach to determine the most likely age–depth relation, resulting in greater flexibility regarding, e.g., the accumulation rates between two dating points. Prior to the actual age modeling, all radiocarbon dates were converted into calendar ages using the *Marine13*-calibration curve as proposed by Reimer et al. [100] with the built-in calibration function of the program.

Using historical, pre-bomb samples, Lloyd et al. [76] estimated a local reservoir correction of 140 ± 35 years for the Disko Bay area. This value has been discussed in more detail in Jackson et al. [45] and widely accepted in previous studies [51, 76, 96, 97] and was, therefore, applied here in the same manner during the calibration.

### Bulk parameters (TOC and sand content)

Freeze-dried and homogenized sediments were taken (5-cm intervals) for total organic carbon (TOC) measurement using a Carbon–Sulfur ELTRA Analyser (CS-800, ELTRA) after removal of carbonates by adding hydrochloric acid (37%, 500 μl). The machine was calibrated with a standard before measurements, and the accuracy of these measurements was controlled by additional standard measurements after every 10 samples, the error of our TOC measurements is at ±0.02%.

### Table 1 Results of the radiocarbon dating on core GeoB19927-3

| Sample ID     | Lab ID     | Depth (cm) | 14C-age (year) | 14C-error (years) | Weight (mg) | Material                        | Min age (years BP) | Max age (years BP) | Mean age (years BP) |
|---------------|------------|------------|----------------|-------------------|-------------|---------------------------------|-------------------|-------------------|-------------------|
| GeoB19927-3_33cm | AWI-1468.1.1 | 33         | 1821           | 176               | 0.69        | Mixed benthic forams            | 557               | 1348              | 970               |
| GeoB19927-3_62cm | AWI-1469.1.1 | 62         | 2072           | 177               | 0.62        | Mixed benthic forams            | 1027              | 1764              | 1408              |
| GeoB19927-3_199cm | Poz-85919   | 199        | 3360           | 30                | 15          | Mollusk shells                  | 2818              | 3190              | 3009              |
| GeoB19927-3_410cm | AWI-1259.1.1 | 410        | 4692           | 197               | 0.8         | Mixed benthic forams            | 4478              | 5122              | 4799              |
| GeoB19927-3_411cm | AWI-1261.1.1 | 411        | 4836           | 192               | 1           | Mixed benthic forams            | 4489              | 5127              | 4807              |
| GeoB19927-3_538cm | Poz-85920   | 538        | 5495           | 35                | 10          | Mollusk shells                  | 5618              | 5901              | 5764              |
| GeoB19927-3_561cm | Poz-85921   | 561        | 5720           | 35                | 12          | Mollusk shells                  | 5803              | 6062              | 5994              |
| GeoB19927-3_607cm | Poz-85924   | 607        | 5885           | 35                | 10          | Mollusk shells                  | 6094              | 6369              | 6222              |
| GeoB19927-3_689cm | Poz-85925   | 689        | 6670           | 50                | 10          | Mollusk shells                  | 6821              | 7219              | 7028              |
| GeoB19927-3_767cm | Poz-85926   | 767        | 7410           | 10                | 10          | Mollusk shells                  | 7584              | 7893              | 7740              |
| GeoB19927-3_790cm | Poz-85927   | 790        | 7640           | 50                | 10          | Mollusk shells                  | 7801              | 8071              | 7938              |
| GeoB19927-3_1000cm | AWI-1260.1.1 | 1000       | 8831           | 205               | 0.8         | Planktonic (*N. pachyderma* sin.) and mixed benthic | 8956              | 9876              | 9371              |

Minimum and maximum ages denote the 95% (2σ) uncertainty of the BACON age model.
Sand content (grain size) measurements were performed at 10–30-cm resolution at the Particle-Size Laboratory at MARUM, University of Bremen with a Beckman Coulter Laser Diffraction Particle Size Analyzer LS 13320. The obtained results provide the particle-size distribution of a sample from 0.04 to 2000 µm, and the sand content class used was between 63 and 2000 µm. The average standard deviation integrated over all size classes (63–2000 µm) is better than ±4 vol% (cf., [6] for a detailed explanation).

Sea ice biomarkers (IP25, HBI III and sterols)

For biomarker analysis, ~4 g of freeze-dried and homogenized sediment (10-cm intervals) was extracted using dichloromethane: methanol (2:1 v/v) as a solvent for ultrasonication (3 × 15 min). Beforehand, 9-octylheptadec-8-ene (9-OHD; 0.1 µg/sample), 7-hexynonadecane (7-HND; 0.076 µg/sample), 5α-androstan-3β-ol (Androstanol; 10.7 µg/sample) and 2,6,10.15,19.23-hexamethyltetrasocene (Squalane; 3.2 µg/sample) were added for biomarker quantification. Sterols and hydrocarbons were separated by open silica (SiO2) column chromatography with n-hexane (5 ml) and ethyl-acetate: n-hexane (9 ml; 2:8 v/v) as eluent. The latter fraction was silylated with 200 µl BSTFA (bis-trimethylsilyl-trifluoroacetamide) (60 °C, 2 h).

The identification of the compounds was carried out with a gas chromatograph (Agilent Technologies GC6850, 30 m DB-1MS column, 0.25 mm id, 0.25 µm film) coupled to an Agilent Technologies 5977 C VL MSD mass selective detector (triple-axis Detector, 70 eV constant ionization potential, Scan 50–550 m/z, 1 scan s⁻¹, ion source temperature 230 °C) for HBI and sterol. GC measurements were carried out with the following temperature program for the hydrocarbons: 60 °C (3 min), 150 °C (15 °C min⁻¹), 320 °C (10 °C min⁻¹), 320 °C (15-min isothermal) for the hydrocarbons and 60 °C (2 min), 150 °C (15 °C min⁻¹), 320 °C (3 °C min⁻¹), 320 °C (20-min isothermal) for the sterols. Helium served as carrier gas (1 ml min⁻¹ constant flow). Specific compound identification was based on the comparison of gas chromatography retention times with those of reference compounds and published mass spectra [7, 16, 19, 125]. For the quantification of IP25 and HBI III (Z and E-isomer) their molecular ion (m/z 350 for IP25 and m/z 346 for HBI III ‘Z and E-isomer’ in relation to the abundant fragment ion m/z 266 of internal standard (7-HND) was used (in selected ion monitoring mode, SIM). The different responses of these ions were balanced by an external calibration curve [36]. For the quantification of the sterols (quantified as trimethylsilyl ethers), the molecular ions m/z 470 for brassicasterol (as 24-methylcholesta-5,22E-dien-3β-ol) and m/z 500 for dinosterol (4α,23,24R-trimethyl-5α-cholest-22E-en-3β-ol) were used in relation to the molecular ion m/z 348 for the internal standard Androstanol. All biomarker concentrations were either normalized to the organic carbon (TOC) content or converted to respective accumulation rates (or flux rates) (See data sets available at https://doi.pangaea.de/10.1594/PANGAEA.911365).

The PIP25 indices were calculated by combining IP25 with different phytoplankton markers for semi-quantitative sea ice reconstruction, according to [89]:

$$P_p IP_{25} = IP_{25} / (IP_{25} + (p \times c))$$

where $p$ is the phytoplankton marker concentration [$p=B$(brassicasterol) or D(dinosterol) or III(HBI III)], and $c$ is a balance factor to compensate for a significant concentration difference between IP25 and phytoplankton marker concentration ($c=$ mean IP25 concentration/mean $p$ concentration). When using HBI III as phytoplankton biomarker and IP25 and HBI III concentrations are similar in magnitude, a balance factor is not needed (i.e., $c=1$). The tri-unsaturated HBI ratio “TR25” was calculated according to [9]:

$$TR_{25} = Z / (Z + E)$$

Results

Core chronology and sedimentation rates

The resulting depth–age model ranges between about 10 ka BP and the present (Fig. 2). The sedimentation rates vary between 32 and 172 cm ka⁻¹. However, mostly high sedimentation rates of 100—150 cm ka⁻¹ occur throughout the majority of the recovered intervals. A step-wise decrease in sedimentation rates to about ~80 cm ka⁻¹ (Fig. 2) is observed at about 8–6 ka BP and to about ~30 cm ka⁻¹ at about 3–1 ka BP, respectively. Excess lead is present down to 4.5 cm core depth, indicating that the core top is of recent age and only experienced minor disturbance during coring (see inset in Fig. 2). Caesium (137Cs) was discovered in the topmost sample at 0–1 cm core depth. Accordingly, we assume that the core top is of near-recent age and used an additional tie point of ~50 years (i.e., 0 a BP) for the core top. The basal age of the core was determined by extrapolation of the age model beyond the lowermost radiocarbon dating to the core base within BACON. Consequently, the age model for the lowermost section (1000–1147 cm/> 9.4 ka) should be regarded with caution, and thus accumulation rates were not calculated. Further details of the age modeling are given in Table 1 and Fig. 2. Mass accumulation rates (g cm⁻² ka⁻¹) were calculated based on sedimentation rates and dry bulk density data, and were finally used to convert biomarker concentrations into flux rates.
Bulk parameters (sand content and TOC)

The sand content (63–2000 µm) displays high-amplitude changes between 0.5 and 63% (vol) in the lowermost part of the core (early Holocene), followed by low and stable values of < 2% in the upper part of the core (7.8 ka to present) (Fig. 3a). The TOC contents range from 0 to 1.5% (Fig. 3b) with minimum but increasing values in the early Holocene (10.4–7.8 ka BP), high values of > 1% in the mid to late Holocene (7.8 ka to present).

The accumulation rates (or fluxes) of TOC range from 0 to 1.3 g cm$^{-2}$ ka$^{-1}$ and show distinctly variable but higher values in the early Holocene (9.3–7.8 ka BP), followed by slightly enhanced values of 0.4–1.3 g cm$^{-2}$ ka$^{-1}$ in the mid Holocene (7.8–3.0 ka BP) with a distinct maximum (~1.3 g cm$^{-2}$ ka$^{-1}$) at ~6 ka BP (Fig. 4b). Between 3.0 and 0 ka BP, the TOC flux decreases, reaching a minimum value of 0.16 g cm$^{-2}$ ka$^{-1}$ at the top of the core.

IP$_{25}$ and other biomarkers

IP$_{25}$ concentrations vary between about 0.5 and 1.5 µg/gTOC in the early Holocene (10.4–7.8 ka BP) with a single distinct maximum of ~3.3 µg/gTOC at 7.9 ka BP, followed by a gradual decrease in the mid Holocene (Fig. 3f). Relatively constant and minimum concentrations of IP$_{25}$ (~0.34 µg/gTOC) are observed in the late Holocene period (3.0–0 ka BP). The phytoplankton biomarkers brassicasterol and dinosterol show a very similar trend in their concentration record with (sub-)millennial-scale fluctuations ranging from 0 to 30 µg/gTOC and 0 to 37 µg/gTOC, respectively (Fig. 3e, d). Intervals of enhanced brassicasterol and dinosterol concentrations are observed from about 9.4 to 8.5 ka BP of ~30 and 37 µg/gTOC and from about 7.4 to 6.3 ka BP of ~19 and 27 µg/gTOC, respectively. In the late Holocene (3.0–0 ka BP), the sterols concentrations are still variable but remain generally low. HBI III concentrations are almost zero before 7.4 ka BP (Fig. 3c), and remain generally low throughout the mid Holocene (mean ~0.4 µg/gTOC), except for a prominent peak of 3.6 µg/gTOC at about 7–6.3 ka BP. In the late Holocene, however, HBI III concentrations increase to ca. 0.8 µg/gTOC, except for two prominent peaks of 2.7 and 2.0 µg/gTOC at about 2.1 and 1.3 ka BP, respectively.

IP$_{25}$ fluxes range from 0 to 1.65 µg cm$^{-2}$ ka$^{-1}$ and are—except for the lowermost part—relatively high in the early Holocene, reaching maximum values at 8.6–8.4 ka BP. In the mid Holocene, the IP$_{25}$ fluxes decrease until 3 ka BP, with some peaks of 1.0, 0.8, 0.9, and 0.4 µg cm$^{-2}$ ka$^{-1}$ at about
7, 6.1, 5.4, and 4.5 ka BP, respectively (Fig. 4f). Minimum IP$_{25}$ flux rates of about 0.14 μg cm$^{-2}$ ka$^{-1}$ are observed in the late Holocene. Flux rates of brassicasterol (Fig. 4e) vary between 0 and 23 μg cm$^{-2}$ ka$^{-1}$ with the absolute maximum at ~ 9.3–8.6 ka. While, flux rates of dinosterol vary between 0 and 19 μg cm$^{-2}$ ka$^{-1}$, and attain a maximum of about 19 μg cm$^{-2}$ ka$^{-1}$ at ~ 6.1 ka BP (Fig. 4d). During the mid Holocene, brassicasterol and dinosterol fluxes show a cyclic variability with a decrease in maximum values towards the top. During the last 3 ka BP, minimum fluxes of brassicasterol and dinosterol between 1.1 and 6.8, and 1.9 and 4.9 μg cm$^{-2}$ ka$^{-1}$, respectively, are typical. In general, the HBI III fluxes display a similar pattern to its concentration record throughout the core (Fig. 4c). Low HBI III fluxes of < 0.1 and between 0.1 and 0.4 μg cm$^{-2}$ ka$^{-1}$ occur in the oldest part > 7.5 and between 6.4 and 2.4 ka BP, respectively. Prominent maxima of 1.96, 1.54, and 1.21 μg cm$^{-2}$ ka$^{-1}$ are recorded at 7.0–6.3, ~ 2.0, and 1.4 ka BP, respectively. Both P$_{BI25}$ and P$_{DI25}$ indices are intermediate to high, varying in the range between 0.3 and 1; whereas P$_{III}$IP$_{25}$ index remained high (~ 0.9) in the early Holocene (Fig. 5a). In the mid to late Holocene, P$_{BI25}$, P$_{DI25}$ and P$_{III}$IP$_{25}$ indices display cyclic variability with minima at 7.0–6.3, 5.8–5.5, 4.5–4.2, 3.7–3.2 and 2.1–1.3 ka BP (Fig. 5a;
Fig. 4  a Bulk accumulation rate and accumulation (flux) rates of (b) total organic carbon (TOC) (g cm$^{-2}$ ka$^{-1}$), c  HBI III (µg cm$^{-2}$ ka$^{-1}$), d  dinosterol (µg cm$^{-2}$ ka$^{-1}$), e brassicasterol (µg cm$^{-2}$ ka$^{-1}$), f sea ice proxy IP$_{25}$ (µg cm$^{-2}$ ka$^{-1}$), g δ$^{18}$O record of the NGRIP ice core from Greenland [124], h Summer insolation at 70° N [68]. All plots are shown versus age in years before present (a BP). WIE$_1$ and WIE$_2$ shown as orange bars, are interpreted as (late) winter-ice-edge (WIE) situations. (cf., [12] and discussion for further explanation). Simultaneous peaks in IP$_{25}$ and phytoplankton; brassicasterol and dinosterol, shown as black downward arrows, are interpreted as (polynya-type) reoccurring ice-edge (IE) situations. Gray box marks the core base where the age model is extrapolated and interpreted with caution. Thus, accumulation rates were not calculated. Black solid triangles mark the AMS $^{14}$C-datings
green dots). HBI TR$_{25}$ ratio (Fig. 5b) is highly variable and ranges from ~ 0.5 to 0.8. The ratio is generally low (~ 0.6) in the early Holocene, except for peaks of about 0.7 at ~ 9.1 and ~ 8.7 ka BP. In the mid to late Holocene, it displays cyclic variability and maxima (> 0.7) at 7.1–6.6, 5.9–5.4, 4.6–4.2, 3.8–3.2, and 2.2–1.3 ka BP, coinciding with minima in P$_{III}$IP$_{25}$ index (Fig. 5b; orange bars). In the last 2.3 ka BP, however, TR$_{25}$ displays the highest concentrations with values of ~ 0.8.

**Discussion**

The biomarker approach presented here allows a comprehensive investigation of paleoceanographic changes in NE Baffin Bay, including variations in sea ice cover, primary production and in combination with other proxies—the variability in strengths of oceanic currents (i.e., WGC, BC, etc.) with a ca 100-year time resolution during the Holocene. The Holocene is characterized by significant changes in oceanic forcing as well as Northern Hemisphere solar insolation [68] and, as a result, affecting the advance and retreat of sea ice cover and ice sheets in the region. Furthermore, the combined use of the different PIP$_{25}$ approaches allowed to distinguish between different seasonal ice-edge situations (cf., [12, 89], see “Biomarker proxies for paleo-environmental reconstruction”). Notably, the phytoplankton biomarkers brassicasterol and dinosterol from our records show a high correlation ($R^2 = 0.8$) suggesting co-production of these sterols in a similar open marine environment. This is also reflected in similar trends in their P$_{BI}IP_{25}$ and P$_{III}IP_{25}$ indices (Fig. 5a). Our combined proxy record based on sea ice (IP$_{25}$), phytoplankton biomarkers (brassicasterol, dinosterol, and HBI III) and bulk parameters indicate a consistent presence of seasonal sea ice (Figs. 3, 6) during the Holocene, although the extent and duration of ice cover situation have changed throughout the Holocene.

**Sea ice variations in the NE Baffin Bay during the Holocene**

**Early Holocene**

From final deglacial cold phase to early Holocene warming between ~ 10.4 (?) and ~ 8.5 ka BP. In the lower-most period (1147–1070 cm / > 9.4 ka; see Supplementary Fig. S1), minimum values to the absence of ice algae biomarkers, phytoplankton biomarkers, TOC and TR$_{25}$ but high P$_{III}IP_{25}$ (~ 0.9) index (Figs. 3, 5) indicate colder conditions with extensive sea ice cover during most part of the year except very short
seasonal break-up periods during late spring and/or autumn (PBIP25 ~ 0.6), coinciding with high ice-rafted debris (IRD) values (indicated by high sand content; Fig. 3a). The high amount of terrigenous material (indicated by high IRD) entrapped during sea ice formation in winter and released nutrients in late spring may have facilitated some primary production (Fig. 3d, e) [106]. The colder conditions in Baffin Bay could also be due to the presence of active ice streams [1, 32] and strongly stratified conditions, possibly related to glacial outwash events and melt-water influx from the retreating ice, and a limited strength of the WGC [52, 70, 101].

Following this initial period, the interval ~9.4–8.5 ka BP (1070–890 cm) is marked by pronounced peaks in the fluxes of IP25, phytoplankton biomarkers brassicasterol and dinosterol and TOC as well as TR25 (Figs. 4, 5), pointing all towards rapid changes in the area characterized by variable/less sea ice to ice-edge (MIZ) conditions and high in situ productivity (Fig. 5a). This reduction in sea ice and increased marine productivity might point towards warmer surface conditions in the early Holocene, marked by strong atmospheric forcing (Fig. 7i). Low values of PBIP25 and PIP25 indices, coinciding with low IRD (Figs. 3a, 5a), may reflect a retreat in sea ice cover (and glaciers) and indicate a phase of more open-water conditions in late spring and/or autumn. Based on the PIIIIP25 values, high sea ice cover might have persisted until early-springs seasons (Fig. 5a). These numbers, however, have to be interpreted very cautiously due to the extremely low HBI III values.

We think this is linked to the northward penetration of the WGC, transporting warm Atlantic Water along the West Greenland coast since at least ~10.4–9 ka BP [41, 52] and, thus, causing higher sea-surface temperatures in northern Baffin Bay. Knudsen et al. [59] reported an increase in calcareous benthic foraminiferal flux and heavier δ18O_benthic values through this interval supporting the influence of WGC up to the northern Baffin Bay and subsequently more open surface waters and reduced sea ice coverage. Based on bowhead bone remains [34], IP25 record from the Canadian Arctic Archipelago Vare et al. [122], and ice-edge indicator foraminifera species (S. feylingi) from Upernavik area Hansen et al. [41], the authors reported reduced to ice-edge sea ice cover, similar to our findings in this study. Based on foraminifera, Ostermann and Nelson [95] reported a similar influx of relatively warm, high saline water into the north of Baffin Bay between ~9.8 and 8.5 ka BP. Furthermore, Levac et al. [71] reported an increase in dinoflagellate abundances from a sediment core in the northern Baffin Bay, suggesting similar warming of surface waters.

The interpretation of continued strong melting with warmer summer waters in Baffin Bay corresponds to the significant climate warming in the early Holocene, widely recognized in the Northern Hemisphere [56, 123]. These warm conditions might have led to high sedimentation rates as a result of increased marine productivity, ice melting, and glacially derived material from adjacent ice sheets in NE Baffin Bay [21, 41]. This is in agreement with high accumulation rates and relatively high terrigenous detritus (Figs. 2, 6).
3a) recorded in the sediments in this interval, which may indicate a strong melt-water input associated with melting of sea ice, GIS and nearby glaciers such as Ussing Braeer and Cornell glacier located near Melville Bugt [54, 56]. Notably, the period with maximum dinosterol flux (Fig. 4d) and spring ice melting in Baffin Bay coincides with maximum Agassiz Ice Cap melt percent (~9.4–8.5 ka BP) [37] and maximum summer insolation at 70° N or higher latitudes ([68] (Fig. 7g, i), which suggests significant warm conditions and increased marine productivity in the area. Ice retreat velocity data (~4.8 km a⁻¹) from Jakobshavns Isbrae also indicate similar rapid deglacial melting following increased atmospheric temperatures at about 9.4–8.5 ka BP ([80, 124]. Additionally, the increase in the amount

**Fig. 7** Comparison of different environmental proxy records around and from Greenland. a PbIP₂₅ record from the Melville Bugt (Core GeoB19927-3, this study), b PbIP₂₅ record from East Greenland Shelf (Core PS2641; [60]), c Fram Strait (Core MSM5/5-712-2; [88]), d dinocyst-based sea ice cover reconstruction (months of sea ice/year) from Disko Bay (Core MSM343300; [96]), e dinocyst-based sea ice cover reconstruction (months of sea ice/year) from Upernavik (Core AMD14-204C; [22]), f diatom-based SST reconstruction from West Greenland (Core MSM343300; [64]), g the Agassiz Melt Layer Record [37], h the δ¹⁸O record of the NGRIP ice core from Greenland [124] and i the summer insolation at 70° N [68]. Black solid triangles mark the AMS ¹⁴C-datings
of pollen grains from Ellesmere Island could also indicate the maximum summer melting between ~9.8 and 8.5 ka BP [17, 66]. This characteristic warming, reduction of spring/summer sea ice and deglacial melting of the GIS have been widely reported in the areas such as Fram Strait and Disko Bay based on PIP25 (Fig. 7c) and SST record (Fig. 7e) [64, 88, 132].

Extended sea ice cover and opening of connection to the Arctic between ~8.5 and ~7.8 ka BP Despite the high solar insolation and deglacial melting having culminated, environmental conditions in northern Baffin Bay remained cold and unstable between ~8.5 and ~7.8 ka BP [35] likely due to counter-effect of the Laurentide, Greenland and Innuitian ice sheets and a connection to the Arctic Ocean [59]. A distinct decrease in ice algae and phytoplankton productivity (Fig. 4d–f), coinciding with peaks in IRD (Fig. 3a) during this interval, indicates a period of major environmental change in the area. Cold surface water conditions and increased, partly almost closed sea ice cover in this period are indicated by the PBIP25, PDIP25, and PIIIIP25 (Fig. 5a). Furthermore, contemporaneous peaks in accumulation rates of IP25 and the phytoplankton biomarkers brassicasterol and dinosterol (Fig. 4d–f) display reoccurring spring ice-edge conditions. This may be related to oscillations in the strengths of the WGC and the melting of sea ice [22, 59, 96]. Interestingly, the HBI III record shows a pronounced maximum between 6.3 and 7.0 ka BP, followed by three smaller peaks of elevated HBI III values at 5.7–5.2, 4.5–4.3, and 3.6–3.3 ka BP (Fig. 4c). According to Belt et al. [12], such HBI III maxima may indicate increased phytoplankton productivity and winter-ice-edge limit (Fig. 6b). Baffin Bay is strongly influenced by seasonal melt-water inflow from the GIS, wind favored deep upwelling and light availability, which may, in turn, also influence the production of HBI III [11, 44]. The strong seasonality in this area may explain such an elevated occurrence of HBI III in ice-edge scenarios. Peaks in HBI III associated with enhanced growth of pytoflagellates near MIZ, may also indicate an up-welling situation (i.e., sites near polynya settings; NWP) caused by wind favored mixing in the late-winter months, when north-westerly wind stress is stronger [26, 59, 71, 84, 86, 119]. The PIIIIP25 indices display minima at 7.0–6.3, 5.8–5.5, 4.5–4.2, and 3.7–3.2 ka BP, coinciding with maxima in the TR25 record, which may be interpreted as ice-edge (polynya-type) situation with reduced sea ice cover and elevated primary production during late winter/early spring (Fig. 5b). The intervals in between are characterized by more extended late winter/early spring sea ice conditions.

Mid Holocene

A cyclic change of minima and maxima in sea ice cover is evident after 7.8 ka BP, as shown by PIP25 index cycllicity (Fig. 5a). Interestingly, the HBI III record shows a pronounced maximum between 6.3 and 7.0 ka BP, followed by three smaller peaks of elevated HBI III values at 5.7–5.2, 4.5–4.3, and 3.6–3.3 ka BP (Fig. 4c). According to Belt et al. [12], such HBI III maxima may indicate increased phytoplankton productivity and winter-ice-edge limit (Fig. 6b). Baffin Bay is strongly influenced by seasonal melt-water inflow from the GIS, wind favored deep upwelling and light availability, which may, in turn, also influence the production of HBI III [11, 44]. The strong seasonality in this area may explain such an elevated occurrence of HBI III in ice-edge scenarios. Peaks in HBI III associated with enhanced growth of pytoflagellates near MIZ, may also indicate an up-welling situation (i.e., sites near polynya settings; NWP) caused by wind favored mixing in the late-winter months, when north-westerly wind stress is stronger [26, 59, 71, 84, 86, 119]. The PIIIIP25 indices display minima at 7.0–6.3, 5.8–5.5, 4.5–4.2, and 3.7–3.2 ka BP, coinciding with maxima in the TR25 record, which may be interpreted as ice-edge (polynya-type) situation with reduced sea ice cover and elevated primary production during late winter/early spring (Fig. 5b). The intervals in between are characterized by more extended late winter/early spring sea ice conditions.
strengthening of WGC and Irminger Current since ~7.8 ka BP [4, 34, 47, 51, 55, 77, 96]. Furthermore, between ~7.3 and ~6.2 ka BP, a reduced sea ice cover and relatively warm WGC were recorded at several sites, including northern Baffin Bay and the eastern Labrador Sea [30, 97]. Thomas et al. [120] have shown a major increase in winter snowfall during the Late Holocene caused by reduced sea ice and open surface water conditions in Baffin Bay and the Labrador Sea, consistent with the occurrence of polynya-type (winter) ice-edge variability. Slightly less pronounced, but warming trend is also observed from the Disko Bay area, based on diatoms-based SST data (Fig. 7f) [64]. Many marine and terrestrial records have also reported the GIS retreat from Disko Bay during this time interval [77, 79, 115], in agreement with our reconstructions in NE Baffin Bay.

Late Holocene

During the last 3 ka BP, the concentrations and accumulation rates of IP25, marine sterols (Figs. 3, 4d–f), and related PIP25 index values remain relatively low and do not reflect the Late Holocene ‘Neoglacial’ cooling trend that follows the decreasing insolation pattern [68] (Fig. 7a, i), widely observed in Northern Hemisphere, such as in eastern Baffin Bay, the Fram Strait and the Canadian Arctic Archipelago area (Fig. 7d, f) [41, 64, 89, 96, 129]. This difference might be due to the dominance of the WGC along the West Greenland coast and sea ice interactions with the adjacent fjord, which may further mitigate sea ice growth. Furthermore, our PIP25 indices show relatively moderate values (0.2–0.6), suggesting a reduced sea ice cover, especially between about 2.4 and 1.0 ka BP (Fig. 5a). Sea ice might have only occurred during the late winter/early spring of this time span, as shown by minimum PIP25 values and the HBI III maxima (Figs. 4c, 5a). Notably, the pronounced peaks in HBI III flux and TR25 (Fig. 5b) at about 2.1 and 1.3 ka BP that might coincide with the Roman Warm Period (RWP) and Medieval Climate Anomaly (MCA), respectively, can be associated with stronger WGC pulses. As our core site is not too distant from NWP, a polynya-type, marginal ice zone situation, driven by northerly winds and occasional upwelling during the warm periods is another option to explain increased productivity and related elevated HBI III values. Similarly, Knudsen et al. [59] reported an increase in large diatoms between 2 and 0.6 ka BP, interpreted as warmer conditions and high productivity in Disko Bay. Based on foraminiferal and sedimentological proxies from West Greenland, Lloyd et al. [78] and Norgaard-Pedersen and Mikkelsen [93] also observed relatively warm oceanic conditions at ~2–1.4 ka BP. Furthermore, based on benthic foraminifera analysis, Perner et al. [97] described a period of warming from ~1.4 to 0.9 ka BP and a relatively warm phase at ~1.8 ka BP and suggested an increased IC contribution to the WGC during the RWP and MCA. Perner et al. [97], Kolling et al. [60] and Allan et al. [2] suggested an enhanced Atlantic Water inflow to the WGC and reduced (winter-only) sea ice between ~1.4–0.9 ka BP and ~1.8 ka BP, with beneficial conditions for phytoplankton blooms, in agreement with our findings.

Several studies have recognized a complex feedback system between Arctic sea ice and the North Atlantic Oscillation (NAO) and reported an anti-phase correlation to the NAO mode associated with the RWP and MCA [63, 67, 99, 108]. However, our records point towards an in-phase response of sea ice cover with changes in NAO modes (i.e., warming during the positive NAO mode associated with the RWP and MCA [74, 121]. However, it should be noted that NAO reconstructions are still subject to debate and far from being fully understood (cf., [62, 108] for a detailed discussion).

Based on an IP25 record from the East Greenland shelf (Fig. 7b), Kolling et al. [60] reported that the PIP25 index therein also does not reflect the ‘Neoglacial’ cooling trend, similar to the records of our study. Based on diatom and foraminifera assemblages, Moros et al. [85] and Caron et al. [22] (Fig. 7e) suggested warm, stratified and highly productive waters in Baffin Bay area [83, 104], in agreement with our interpretation. Glaciers, land-fast ice, and local fjords may also have a more direct and unfavorable influence on the growth of sea ice cover [103].

Summary and conclusions

Holocene sea ice conditions and marine phytoplankton productivity were reconstructed to gain direct insights into the sea ice variability and its driving mechanisms using a multi-proxy biomarker approach. Organic geochemical and biomarker analyses of a well 14C-AMS- and 210Pb-dated sediment core from NE Baffin Bay show that major environmental and paleoceanographic changes occurred in this area.

In the lower-most part, a cold interval characterized by extensive sea ice cover and very low local productivity is succeeded by an interval (~9.4–8.5 ka BP) of persistent, albeit strongly variable (reduced) sea ice cover, enhanced GIS spring melting, and strong influence of the WGC in the earliest part of the record. A short-term cooling event is recorded by the ice algae and phytoplankton biomarkers between 8.5 and 7.8 ka BP, pointing towards an increased seasonal sea ice cover as a result of the opening of Nares Strait which led to an increased influx of Polar Water into the Baffin Bay, albeit insolation remained generally high.

The interval between 7.8 and 3.0 ka BP is characterized by reduced sea ice with millennial-scale variability of (late winter) ice-edge limit and WGC strength, and increased open-water (polynya-type) conditions corresponding to ‘warmer’ conditions.
Our IP$_{25}$-based sea ice reconstructions and related PIP$_{25}$ index do not reflect the late Holocene Neoglacial cooling trend during the last 3 ka BP, probably due to the strong influence of the WGC and interactions with the adjacent fjords. Peaks in HBI III at about 2.1 and 1.3 ka BP might coincide with the RWP and MCA, respectively, and are associated with an enhanced WGC and in-phase correlation with NAO mode.

The result of this multi-proxy approach presented here seems to display the rapid transitions between different climate events and indicate a close connection of sea ice variations in the Northern Hemisphere driven by the interplay between melt-water discharge and solar and oceanic forcings (i.e., WGC, BC). These findings may help to understand the recent reduction in sea ice cover and further improve our climate models and climate predictions. More high-resolution studies are needed to understand the complex interaction of sea ice, its driving mechanisms, and fjord interactions in Baffin Bay.

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Compliance with ethical standards

Conflict of interest On behalf of all authors, the corresponding author states that there is no conflict of interest.

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