Changing sources and burial of organic carbon in the Chukchi Sea sediments with retreating sea ice over recent centuries

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**Abstract.** Decreasing sea ice extent caused by climate change is affecting the carbon cycle of the Arctic Ocean. In this study, surface sediments across the western Arctic Ocean are investigated to characterize sources of sedimentary organic carbon (OC). Bulk organic parameters (total organic carbon, total nitrogen, $\delta^{13}$C$_{org}$ and $\delta^{15}$N) and molecular organic biomarkers (e.g., sterols and highly branched isoprenoids (HBIs)) are combined to distinguish between sympagic, pelagic, and terrestrial OC sources. Their downcore profiles generated at the Chukchi Sea R1 core site (74°N) are then used to evaluate changes in the relative contribution of these components of sedimentary OC over the last 200 years with decreasing sea ice. Our data evidence that from 1820s to 1930s prevailing high sea ice cover inhibited *in situ* primary production resulting in prominent land-derived material in sediments. Then, from 1930s to 1980s, primary production starts increasing with the gradual decline of summer sea ice. The ratio of sympagic and pelagic OC began to rise to account for the larger portion of sedimentary OC. Since 1980s, accelerated sea ice loss led to enhanced primary production, stabilizing over the last decades due to freshwater induced surface ocean stratification in summer.
1 Introduction

Knowledge on processes and feedback mechanisms controlling the carbon cycle is essential for a better understanding of Arctic marine ecosystem dynamics and its role in climate change (Parmentier et al., 2017; Wheeler et al., 1996). The Arctic Ocean is the major world carbon sink region where huge amounts of marine and terrestrial organic carbon (OC) have accumulated (Stein et al., 2004). Today, the Arctic Ocean experiences unprecedented changes caused by global warming and Arctic amplification (Cavalieri et al., 1997; Rantanen et al., 2022; Serreze and Francis, 2006; Shindell and Faluvegi, 2009) which have resulted in major sea ice loss with consequences on the marine ecosystems and Arctic carbon budget. Increased river discharge and melting permafrost are responsible for enhanced delivery of terrigenous inorganic and organic carbon to the Arctic marginal seas (Grotheer et al., 2020; Holmes et al., 2011; Rawlins et al., 2021; Vonk et al., 2012). Terrigenous OC reaching the Arctic Ocean is either partly mineralized or transported to the sea floor where it is ultimately buried (Fritz et al., 2017; Tanski et al., 2019). Increased nutrient-rich waters brought by enhanced Pacific water inflow (PWI), a major source of nutrients to the Arctic, also contribute to stimulate phytoplankton productivity in the western Arctic Ocean (Arrigo and van Dijken, 2015; Tian et al., 2021; Woodgate and Peralta-Ferriz, 2021; Woodgate, 2018). Lastly and most importantly, the rapid sea ice loss in summer has resulted in large areas of the Arctic Ocean that shifted from multi-years to seasonal sea ice coverage (Cavalieri and Parkinson, 2012; Parkinson et al., 1999; Stroeve et al., 2007) allowing higher light penetration in surface waters thereby enhancing primary production and export of OC to the bottom floor. Enhanced summer sea ice melting further contributes to sea ice algal production, export and burial of marine OC in sediments (Ardyna and Arrigo, 2020). The Chukchi Sea (CS) is one of the most productive regions of the Arctic marginal seas (Cai et al., 2010; Ouyang et al., 2022; Zhuang et al., 2022). With the rapid sea ice retreat, the CS has become a key area to study climate induced OC changes since the beginning of the Industrial Era.

A large variety of indicators, including bulk geochemical ratios and lipid biomarkers (Volkman, 1986) have been developed to characterize the composition of OC in Arctic Ocean sediments (Fernandes and Sicre, 2000; Sparkes et al., 2015). Among them, lignins and δ^{13}C_{org} have been successfully used to provide reliable estimates of terrestrial OC (Tesi et al., 2014; Wang et al., 2019a; Wild et al., 2022).
However, pelagic and sympagic sourced OC remains difficult to discriminate in the Arctic Ocean, particularly in regions of high sympagic productivity. To address this issue, the H-print index based on highly-branched isoprenoids (HBIs) defined as the ratio of pelagic HBI-III over the sum of HBIs (IP$_{25}$ + HBI-II + HBI-III) was developed (Brown et al., 2014b; Koch et al., 2020). Values close to 100% are thus indicative of prominent pelagic sources while those close to 0% reflect prevailing sympagic sources. By combining H-print and $\delta^{13}$C$_{org}$, we here intend to more accurately quantify marine pelagic, marine sympagic and terrestrial fractions of OC in Arctic sediments.

Documenting changes in sea ice changes and induced transformations of Arctic ecosystems is key to better predict how the carbon cycle will respond to future changes with continued warming (Arrigo et al., 2008; Bates and Mathis 2009). Extensive surveys of Arctic sea ice have been possible only since the 1970s with the development of remote sensing observations (Cavaliere et al., 1996). Prior to this, very few in situ observations on sea ice exists due to the inaccessibility of the Arctic Ocean. Paleoclimate proxies such as fossils assemblages and geochemical indicators have thus been used as alternative approaches to document past changes of sea ice and place them in the context of ongoing changes (e.g. Belt, 2018; de Vernal et al., 2013 and references therein). The monounsaturated HBI biomarker IP$_{25}$ (Ice Proxy with 25 carbon atoms) produced by sea ice diatoms was initially proposed to assess seasonal sea ice cover (Belt et al., 2007; Massé et al., 2008). The PIP$_{25}$ (Phytoplankton-IP$_{25}$) index, that combines IP$_{25}$ with pelagic phytoplankton biomarkers, was then proposed to provide semi-quantitative estimates of seasonal sea ice (Belt, 2019; Müller et al., 2011). Most IP$_{25}$-related studies in the Arctic Ocean have focused on surface sediments to derive spatial seasonal sea ice distribution (Kolling et al., 2020; Stoynova et al., 2013; Su et al., 2022; Xiao et al., 2015a; Xiao et al., 2013). Others have investigated its past variability at the millennial scale and beyond (Cronin et al., 2013; Polyak et al., 2016; Stein et al., 2017; Xiao et al., 2015b), but only a limited number has explored sea ice variability over the past centuries (Bai et al., 2022; Hu et al., 2020; Kim et al., 2019). None have attempted to link seasonal sea ice changes to sedimentary OC composition since the beginning of the Industrial Era.

In this study, we investigate the potential of H-print combined with $\delta^{13}$C$_{org}$ in surface sediments of the northern CS to discriminate and quantify the relative contribution of OC originating from pelagic, sympagic and terrestrial sources and their evolution over the last two centuries under changing sea ice
conditions to improve our understanding of ongoing alteration of the OC cycle.

Figure 1. Map of the western Arctic Ocean (AO) showing surface ocean circulation and sampling locations (black, gray, and green dots). The stations in gray and green represent the surface samples collected on the LV77 and 11th CHINARE cruises, respectively. The black dot indicates the sediment core ARC11-R1 location. The black diamonds show two other sediment cores discussed in the text (ARC04-C07, Bai et al 2022; ARA01-03MUC, Kim et al 2019). Surface sediments reported by Bai et al. (2019) and Wang et al. (2017) are shown by red and orange dots, respectively. The red start shows the location of the sediment trap station DM (Bai et al., 2019). The dotted and dashed lines in black represent the 20% isolines of September sea ice concentration for the 2019 and 1979, respectively (https://nsidc.org; Cavalieri et al., 1996). Main study regions: ESS, East Siberian Sea; CS, Chukchi Sea; CP, Chukchi Plateau. Main surface currents: SCC, Siberian coastal current; PWI, Pacific water inflow; ACW-Alaskan Coastal Water; AW-Anadyr Water; BSW-Bering Shelf Water. Rivers are shown in green lines: IR, Indigirka River; KR, Kolyma River.

2 Oceanographic setting

The CS is one of the largest marginal seas in the world located on the northern Asian and American continents (Jakobsson, 2002). The surface ocean circulation in the CS is controlled by winds and sea ice cover (Ovall et al., 2021). This basin is connected to the Pacific Ocean through the Bering Strait. The PWI entering the Arctic Ocean strongly influences the physico-chemical water properties of the Arctic Ocean and contributes to enhanced primary production (Coachman and Aagaard, 1966) (Fig. 1). In the CS, the PWI divides into three branches: the highly saline and high-nutrient content Anadyr Water (AW)
on the western side, the fresher and oligotrophic Alaska Coastal Water (ACW) on the eastern side, and the moderately saline Bering Shelf Water (BSW) in between (Grebmeier et al., 2006; Hunt et al., 2013; Woodgate et al., 2005). The eastern side of the CS is adjacent to the Beaufort Sea. The dynamics of the Beaufort Gyre (BG) also impacts on the characteristics of the CS water mass (Timmermans and Toole, 2023). In particular, enhanced anticyclonic BG circulation has resulted in increased freshwater convergence into the Canadian Basin in recent years (Giles et al., 2012) with implications on the local biological production as well as on the transport of terrestrial organic matter (He et al., 2012; Coupel et al., 2015; Ren et al., 2020). Fresh and cold waters from the seasonal Siberian coastal current (SCC) is another feature of the surface ocean circulation that lowers salinity of the central CS water (Weingartner et al., 1999). The dramatic loss of sea ice in summer caused by global warming is particularly tangible in the Arctic Ocean marginal seas (Cavaliere et al., 1997; Parkinson et al., 1999; Polyakov et al., 2003; Zhang et al., 2021) and most pronounced in the CS both in terms of sea ice extent and thickness (Serreze and Stroeve, 2015; Wang et al., 2019b). Remote sensing data (1979 to 2020) reveal considerable seasonal variations of sea ice extent in the CS. The CS is heavily covered by sea ice from November to June. Sea ice gradually decreases in July and reaches its minimum extent in September (https://nsidc.org; Cavaliere et al., 1996).

3 Material and methods

3.1 Material

A total of 42 surface sediments (0-2 cm) from the East Siberian Sea (ESS) and CS were collected during the Cruise LV77 on board the R/V Akademik M.A. Lavrentiev. Additional 11 surface sediments (0-2 cm) and a 15 cm long sediment core ARC11-R1 (R1 hereafter, 74.64°N, 169.13°W, 200 m water depth) were also collected using a box corer and a multi-corer, respectively, in the CS and Chukchi Plateau (CP) during the 11th Chinese National Arctic Research Expedition (CHINARE) in summer 2020 on board the R/V Xuelong 2 (Fig. 1). Subsampling was performed on board at a sampling interval of 1 cm. Subsampled core sediments and surface sediments were frozen immediately after recovery at -20°C until further analysis in the laboratory.

3.2 Sediment core chronology

The chronology of the R1 core is based on excess $^{210}\text{Pb}$ ($^{210}\text{Pb}^{\text{ex}}$) determinations performed at the
State Key Laboratory of Estuarine and Coastal Research, East China Normal University, Shanghai, China, using an HPGe gamma spectrometry (GSW275L, Canberra). The $^{210}$Pb$_{ex}$ activity was calculated by subtracting the supporting fraction ($^{226}$Ra) from the total $^{210}$Pb ($^{210}$Pb$_{total}$) activity in the sediment. The error in $^{210}$Pb$_{ex}$ is computed by propagating the error in the corresponding measured pair ($^{210}$Pb and $^{226}$Ra).

A mean linear sedimentation rate (cm yr$^{-1}$) was calculated from the $^{210}$Pb$_{ex}$ profile using a Constant Flux-Constant Sedimentation Rate (CF-CS) model, assuming continuous homogeneous deposition of non-equilibrium $^{210}$Pb in the sediment (Nittrouer et al., 1984). As $^{137}$Cs was also measured during the same gamma counting session, this radionucleide was also used to test the chronology.

### 3.3 Bulk analyses

Total organic carbon (TOC), total nitrogen (TN), $\delta^{13}$C$_{org}$, and $\delta^{15}$N of the 42 surface sediments from the ESS and western CS during the LV77 cruise as well as the 11 surface sediments and the R1 core retrieved from the CS during the CHINARE cruise were analyzed at the Key Laboratory of Marine Ecosystem Dynamics, Second Institute of Oceanography, Ministry of Natural Resources (MED, SIO, MNR, Hangzhou, China). There were first freeze-dried, then ground and homogenized before analyses. About 0.5 g of sediment was acidified using 1 mol L$^{-1}$ HCl and heated overnight in a water bath at 50°C. The excess acid was washed away using ultrapure water (Williford et al., 2007). These samples were weighed for TOC and $\delta^{13}$C$_{org}$ determinations. For TN and nitrogen stable isotopes ($\delta^{15}$N) analyses, we used samples that were not acidified. TOC, TN, $\delta^{13}$C$_{org}$, and $\delta^{15}$N measurements were carried out on an elemental analyzer Elementar CHNOS coupled to an isotope ratio mass spectrometer (IRMS, Thermo, Delta V advantage). The standard deviations for TOC, TN, $\delta^{13}$C$_{org}$, and $\delta^{15}$N based on replicate analyses were 0.02%, 0.005%, 0.2‰, and 0.2‰, respectively.

### 3.4 Biomarker Analyses

Biomarker analyses were completed at MED, SIO, MNR (Hangzhou, China). Before extraction, internal standards 7-hexynonadecane and cholest-5-en-3β-ol-D6 were added to about 5 g of freeze-dried and homogenized sediment for quantification of HBIs and sterols, respectively. Extraction was performed 3 times in an ultrasonic bath for 15 min using dichloromethane/methanol (2:1 v/v). The 3 extracts were combined and dried under a gentle nitrogen stream. Further separation was carried out by adsorption chromatography on an open-column filled with SiO$_2$ using 2.5 ml n-hexane and 4 ml n-
hexane/ethyl acetate (70:30 v/v) to separate the hydrocarbons and sterols, respectively, from the total lipid extract. About 50 μl BSTFA (bis-trimethylsilyl-trifluoroacetamide) were added to the sterol fraction and heated at 70° C for 1 hr for silylation.

Then, hydrocarbons and sterols were analyzed by gas chromatography (GC, Agilent Technologies 7890, 30 m HP-1MS column, 0.25 mm in diameter, and 0.25 μm film thickness) coupled to mass spectrometry (MS, Agilent 262 Technologies 5975C inert XL). A heating rate of 10°C min⁻¹ for the oven temperature was programmed from 40°C to 300°C and maintained at final temperature for 10 min. The ion source temperature was set at 250°C and ionization energy at 70 eV for MS analyses (Belt et al., 2007, Müller et al., 2009). Individual compounds were identified based on their retention time and mass spectra. Selective ion monitoring was used to detect the C_{25}-HBIs (m/z 350 for IP_{25}, m/z 348 for HBI-II, and m/z 346 for HBI-III) and the sterols (m/z 470 for brassicasterol (24-methylcholesta-5,22E-dien-3β-ol), m/z 500 for dinosterol (4α,23,24R-trimethyl-5α-cholest-22E-en-3β-ol), m/z 396 for β-sitosterol (24-ethylcholesterol-5-3β-ol) and m/z 382 for campesterol (24-methylcholesterol-5-3β-ol). Concentrations of HBIs were determined based on the area of individual compounds and that of the 7-hexylnonadecane (m/z 266) obtained by GC/MS. Similarly, sterol concentrations were calculated from the area of individual sterols and cholesterol-D6 (cholesterol-5-en-3β-ol-D6, m/z 464) (Belt et al., 2012, Müller et al., 2011). Concentrations of all biomarkers were normalized to TOC.

### 3.5 PIP_{25} Index and H-print

PIP_{25} indexes were calculated to estimate seasonal sea ice concentrations (Müller et al., 2011) using the following expression:

\[
\text{PIP}_{25} = \frac{[\text{IP}_{25}]}{[\text{IP}_{25}] + [\text{phytoplankton biomarker}] \cdot c}
\]

where

\[
\begin{align*}
\text{c} &= \frac{\text{mean } \text{IP}_{25} \text{ concentration}}{\text{mean } \text{phytoplankton biomarker concentration}} \\
\end{align*}
\]

Brassicasterol (B), dinosterol (D), and HBI-III (III) were used as a reference for pelagic phytoplankton to calculate the P_{B}IP_{25}, P_{D}IP_{25}, and P_{III}IP_{25} values, respectively.

The H-print values were also calculated to infer the relative contribution of pelagic and sympagic OC (Brown et al., 2014b, Koch et al., 2020).
\[ H - \text{print\%} = \frac{[HBI - III]}{[IP_{25}] + [HBI - II] + [HBI - III]} \times 100 \] \hspace{1cm} (3)

Low H-print values are indicative of higher sympagic production while high H-print values point to prevalent pelagic algae.

3.6 Environmental dataset of surface sediment

Assessment of seasonal sea ice spatial distribution is based on the compilation of previously published surface sediment data from the ESS (HBIs, Su et al., 2022) and the CS (HBIs, Bai et al., 2019; \(\delta^{13}C_{org}\), Wang et al., 2017) and new data from the CS and CP produced in this study. \(\delta^{13}C_{org}\) data from the ESS are also new together with all data from the R1 core.

4. Results

4.1 Chronology of core R1

![Figure 2](image)

The activity of \(^{210}\text{Pb}_{\text{total}}\) ranges from 22 to 214 Bq Kg\(^{-1}\), with an average value of 64.30 Bq Kg\(^{-1}\) (Fig. 2). \(^{210}\text{Pb}_{\text{ex}}\) decreases exponentially with increasing depth to reach negligible values around 9 cm. The calculated average sedimentation rate of R1 using the CF-CS model is estimated to 0.07 cm yr\(^{-1}\) \((r^2=0.94, p<0.05)\), which gives a time span of about 200 years for the whole core. The \(^{137}\text{Cs}\) profile (not shown)
### Table 1. Summary of bulk parameters and biomarker data from surface sediment in the Chukchi Sea and Chukchi Plateau.

<table>
<thead>
<tr>
<th>Station</th>
<th>Long.</th>
<th>Lat.</th>
<th>TOC (wt%)</th>
<th>TN (wt%)</th>
<th>(\delta^{13}C_{org}) (%)</th>
<th>(\delta^{15}N) (%)</th>
<th>IP(_{25}) relative abundance(^*)</th>
<th>HBI-II relative abundance(^*)</th>
<th>HBI-III relative abundance(^*)</th>
<th>Brassinosterol(^*)</th>
<th>Dinosterol(^*)</th>
<th>Campesterol+(\beta)-sitosterol(^*)</th>
</tr>
</thead>
<tbody>
<tr>
<td>E1</td>
<td>-179.89</td>
<td>75.01</td>
<td>1.23</td>
<td>0.17</td>
<td>-23.72</td>
<td>9.42</td>
<td>1.58</td>
<td>1.23</td>
<td>0.17</td>
<td>25.34</td>
<td>3.11</td>
<td>189.29</td>
</tr>
<tr>
<td>P2-5</td>
<td>-163.68</td>
<td>76.60</td>
<td>0.87</td>
<td>0.12</td>
<td>-22.73</td>
<td>8.67</td>
<td>0.22</td>
<td>0.18</td>
<td>0.04</td>
<td>1.79</td>
<td>0.60</td>
<td>37.33</td>
</tr>
<tr>
<td>R5</td>
<td>-168.94</td>
<td>77.76</td>
<td>1.18</td>
<td>0.15</td>
<td>-23.54</td>
<td>9.05</td>
<td>0.52</td>
<td>n.d.</td>
<td>n.d.</td>
<td>3.05</td>
<td>0.92</td>
<td>44.19</td>
</tr>
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<td>E2</td>
<td>179.99</td>
<td>75.84</td>
<td>0.57</td>
<td>0.13</td>
<td>-22.99</td>
<td>8.92</td>
<td>3.90</td>
<td>n.d.</td>
<td>n.d.</td>
<td>20.07</td>
<td>3.59</td>
<td>127.45</td>
</tr>
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<td>R2</td>
<td>-168.92</td>
<td>75.61</td>
<td>0.96</td>
<td>0.18</td>
<td>-22.64</td>
<td>9.91</td>
<td>1.00</td>
<td>0.69</td>
<td>0.76</td>
<td>19.30</td>
<td>4.28</td>
<td>135.13</td>
</tr>
<tr>
<td>P3-7</td>
<td>-165.92</td>
<td>78.61</td>
<td>0.65</td>
<td>0.12</td>
<td>-22.26</td>
<td>8.01</td>
<td>n.d.</td>
<td>n.d.</td>
<td>n.d.</td>
<td>3.27</td>
<td>0.65</td>
<td>63.21</td>
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<td>P1-6</td>
<td>-166.62</td>
<td>75.44</td>
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<td>0.16</td>
<td>-23.13</td>
<td>9.44</td>
<td>1.00</td>
<td>0.59</td>
<td>1.47</td>
<td>7.85</td>
<td>2.81</td>
<td>86.21</td>
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<td>Z4</td>
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<td>73.54</td>
<td>1.42</td>
<td>0.28</td>
<td>-22.89</td>
<td>8.74</td>
<td>5.49</td>
<td>6.52</td>
<td>3.06</td>
<td>21.89</td>
<td>7.04</td>
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<td>n.d.</td>
<td>n.d.</td>
<td>n.d.</td>
<td>5.56</td>
<td>1.02</td>
<td>79.41</td>
</tr>
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</table>

n.d.: not detected

\(\delta\) in \(\mu g g^{-1}\) TOC

### Table 2. Summary of bulk parameters and biomarker data from core ARC11-R1.

<table>
<thead>
<tr>
<th>Core depth (yr AD)</th>
<th>Age</th>
<th>TOC (wt%)</th>
<th>TN (wt%)</th>
<th>C/N Ratio</th>
<th>(\delta^{13}C_{org}) (%)</th>
<th>(\delta^{15}N) (%)</th>
<th>IP(_{25}) relative abundance(^*)</th>
<th>HBI-II relative abundance(^*)</th>
<th>HBI-III relative abundance(^*)</th>
<th>Brassinosterol(^*)</th>
<th>Dinosterol(^*)</th>
<th>Campesterol+(\beta)-sitosterol(^*)</th>
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<td>0-1</td>
<td>2013</td>
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<td>0.18</td>
<td>7.76</td>
<td>-23.51</td>
<td>9.35</td>
<td>0.45</td>
<td>0.77</td>
<td>0.51</td>
<td>40.04</td>
<td>8.73</td>
<td>86.00</td>
</tr>
<tr>
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<td>1.13</td>
<td>0.17</td>
<td>7.66</td>
<td>-23.79</td>
<td>9.32</td>
<td>0.66</td>
<td>0.88</td>
<td>0.62</td>
<td>34.25</td>
<td>5.63</td>
<td>69.22</td>
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<td>0.15</td>
<td>7.15</td>
<td>-23.46</td>
<td>9.05</td>
<td>0.82</td>
<td>0.93</td>
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<td>5.09</td>
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<td>0.11</td>
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<td>7.53</td>
<td>0.24</td>
<td>0.43</td>
<td>0.16</td>
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<td>0.85</td>
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<td>6.56</td>
<td>0.39</td>
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<td>0.10</td>
<td>9.49</td>
<td>-24.63</td>
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\(\delta\) in \(\mu g g^{-1}\) TOC
further supports the $^{210}\text{Pb}$ dating. This value falls within the range reported by Cooper and Grebmeier (2018) in a Chukchi Shelf core (0.03-0.37 cm yr$^{-1}$) and is slightly lower than found at the ARC4-C07 core (0.09 cm yr$^{-1}$, Bai et al., 2022) and ARA01B-03MUC core (0.09 cm yr$^{-1}$, Kim et al., 2019) both located to the south (Fig. 1).

![Figure 3](image)

**Figure 3.** Downcore profiles of (a) Total Organic Carbon, TOC in %, (b) Organic carbon isotopic composition ($\delta^{13}\text{C}_{\text{org}}$) of the TOC in ‰, (c) Total Nitrogen, TN in %, (d) nitrogen isotopic composition ($\delta^{15}\text{N}$) in ‰, and (e) C/N ratio in the ARC11-R1 core.

4.2 Proxy data

4.2.1 Surface sediments
The TOC and TN of surface sediments range from 0.57% to 1.42% and from 0.12% to 0.28% respectively (Table 1, Fig. S1). \( \delta^{13}C_{org} \) varies from -23.7‰ to -22.0‰ and \( \delta^{15}N \) from 8.01‰ to 9.91‰.

Both HBIs and pelagic phytosterol concentrations showed a gradual decrease from the shelf to the northern CP. The concentrations of HBI-II and HBI-III reached their detection limit at around 76° N, whereas for IP25 this limit is achieved north of 78° N. By contrast, brassicasterol and dinosterol were detected in all samples with highest values recorded at the shelf edge. Terrestrial sterols (β-sitosterol and campesterol) showed high values over the shelf and minimum ones at the northern end of the CP.

Figure 4. Downcore profiles of (a) IP25, (b) HBI-II, (c) HBI-III, (d) brassicasterol and dinosterol, and (e) campesterol and β-sitosterol in core ARC11-R1.
4.2.2 ARC11-R1 core

The TOC and TN downcore profiles over the last 200 years both show increasing trends towards present ($r^2=0.88$, $p<0.01$) with values varying from 0.71% to 1.18% and from 0.09% to 0.18%, respectively (Table 2, Fig. 3a and c). TOC exhibits a minimum end of the 1950s and rapidly increases thereafter. Downcore values of C/N ratios show a gradual decrease from 9.2 to 7.6 while $\delta^{13}$C$_{org}$ consistently increases from -25.07‰ to -23.46‰ in the 1980s (Table 2, Fig. 3b and d). $\delta^{15}$N exhibits constant values until the early 1900s after which it gradually increases and reaches a maximum at the top of the core (9.35‰; Table 2, Fig. 3d).

The IP$_{25}$ concentrations span from 0.18 to 0.85 $\mu$g g$^{-1}$ TOC with highest values found in the 1930s and after the 1980s (Fig. 4a). Brassicasterol and dinosterol both exhibit increasing abundances over time (Fig. 4d, $r^2=0.85$, $p<0.01$), with brassicasterol being notably more abundant (6.51 $\mu$g g$^{-1}$ TOC to 40.04 $\mu$g g$^{-1}$ TOC) than dinosterol (2.40 $\mu$g g$^{-1}$ TOC to 8.73 $\mu$g g$^{-1}$ TOC; Fig. 4d). Finally, terrestrial sterols slowly increase from 1820s to present in R1 core (campesterol: 10.47 $\mu$g g$^{-1}$ TOC to 68.98 $\mu$g g$^{-1}$ TOC, $\beta$-sitosterol: 4.15 $\mu$g g$^{-1}$ TOC to 17.02 $\mu$g g$^{-1}$ TOC; Fig. 4e).

5 Discussion

5.1 Reconstruction of sea ice conditions

Sympagic biomarker IP$_{25}$ concentrations (0.40 ± 0.25 $\mu$g g$^{-1}$ TOC, mean ± S.D.) were lower throughout the R1 core than found over the same period in the ARA01B-03MUC (0.96 ± 0.72 $\mu$g g$^{-1}$ TOC, Kim et al., 2019) and ARC04-C07 cores (1.29 ± 1.19 $\mu$g g$^{-1}$ TOC, Bai et al., 2022), both located south of R1 core (Fig. S2). Decreasing sympagic biomarker concentrations with increasing latitude likely reflect lower export of sympagic OC to the sea floor due to higher sea ice cover in the North. However, this interpretation does not rule out a possible contribution of analytical methods and/or regional depositional conditions (Belt, 2018). The box-plot in Fig. S2 also points to higher variability at the two further south sites possibly reflecting sea ice edge variations. The presence of IP$_{25}$ throughout R1 indicates seasonal sea ice cover and/or sea ice edge conditions since the 1820s at this location. While Bai et al. (2022) reported parallel trends of IP$_{25}$ and HBI-II throughout core ARC4-C07, this feature is only observed after the 1930s in core R1 (Fig. 4a and b). Indeed, before 1930s, these two HBIs show opposite behavior. The reasons for this discrepancy are not entirely clear and few data exist to explore in depth.
possible explanations. It is worthy to note that both HBIs in the sediment trap at the DM station (74°N) shown similar production/export behavior (Bai et al., 2019). HBI-II is commonly found in Southern Ocean sediments, unlike IP25, where its production has been attributed to the sea ice diatom Berkelya adeliensis (Belt et al., 2016; Brown et al., 2014a). It was also noted that B. adeliensis tends to preferentially flourish in platelet ice, particularly in coastal settings, leading to link its occurrence to landfast sea ice associated with freshwater discharge in Southern Ocean sediments (Belt, 2018, 2019).

The divergent behavior the HBI-II and IP25 prior the 1930s could thus indicate variable sea ice conditions with a possible contribution of drifting ice from coastal areas of the ESS.

Combined IP25 and pelagic phytoplankton biomarkers were investigated to quantify downcore seasonal sea ice cover. To test the sensitivity of PIP25 to c-factor, PIP25 values were calculated using different c-factors of Xiao et al. (2015a) and Smik et al. (2016), and c-factor calculated from surface sediments collected from the CS and R1 for this study. The estimated PIP25 values were found to be consistent with each other with comparable fluctuations (Fig. S3). Additionally, a previous study in the same region suggested that the PIP25 derived sea ice reconstructions were more reliable by using c-factors from the Pan-Arctic database (Kim et al., 2019). Thus, in this study, PbIP25, PdIP25, and PIIIIP25 were calculated using surface sediment balance factors c of 0.02, 0.11 (Xiao et al., 2015a) and 0.63 (Smik et al., 2016), respectively. All indexes show similar trends (Fig. S3) reflecting the strong correlation between brassicasterol, dinosterol and HBI-III (Fig. 5 and S4; all: r>0.63, p<0.01). Only in the 1820s and 1930s are PIP25 values above the threshold of 0.75 indicative of permanent sea ice according to Müller et al. (2011) (Fig. 5).

Between the 1820s and 1850s, all PIP25 values steeply drop suggesting rapid sea ice retreat. Then, PbIP25 and PdIP25 show rather stable values till the beginning of the 20th century whereas PbIIIIP25 slowly increase (Fig. 5). All three PIP25 indexes point to seasonal sea ice or marginal ice zone conditions. Higher amounts of HBI-III and PIIIIP25 values may indicate sea ice edge conditions. Sediment trap data from the Northwind Ridge and slope of the East Siberian Sea show high levels of brassicasterol till late summer/early autumn when ice free conditions are reached, whereas HBI-III is close to the detection limit in mid-summer when sea ice has melted (Bai et al., 2019, Gal et al., 2022). This result further confirms that HBI-III producers proliferate at the sea ice edge rather than in ice free waters. From 1850s
to 1910s, both P\textsubscript{III}IP\textsubscript{25} and P\textsubscript{III}IP\textsubscript{25} were below 0.5, pointing to less and variable sea ice cover than the end of early 19th century, contrary to observations indicating high sea ice cover (Walsh et al., 2017). P\textsubscript{III}IP\textsubscript{25} falls in a higher sea ice cover range. Increasing P\textsubscript{III}IP\textsubscript{25} and P\textsubscript{III}IP\textsubscript{25} values between 1910s and 1930s occur while observations indicate a significant reduction of sea ice from permanent to marginal sea ice (Walsh et al., 2017). Bias to higher sea ice cover estimates using P\textsubscript{III}IP\textsubscript{25} is explained by HBI-III production taking place at the ice edge rather than ice-free conditions. Low concentrations of IP\textsubscript{25} and phytoplankton biomarkers during this period may also uncertainties in the calculation of the PIP\textsubscript{25} index (Fig. 6) as suggested by previous studies (Müller et al., 2011; Xiao et al., 2015b). Observation data of September sea ice extent since 1850 in the Chukchi Sea (Walsh et al., 2017) also supports high sea ice conditions between 1820s and 1930s (Fig. 5).

**Figure 5.** PIP\textsubscript{25} index using brassicasterol (in green, P\textsubscript{III}IP\textsubscript{25}, c = 0.02 as calculated by Xiao et al. (2015a)), dinosterol (in purple, P\textsubscript{III}IP\textsubscript{25}, c = 0.11 as calculated by Xiao et al. (2015a)), and HBI-III (in orange, P\textsubscript{III}IP\textsubscript{25}, c = 0.63 as calculated by Smik et al (2016)) and September sea ice extent record for the Chukchi Sea (red line, Walsh et al., 2017).

**Figure 6.** Cross-plot between concentrations of IP\textsubscript{25} and phytoplankton biomarkers ((a) HBI-III, (b) brassicasterol and (c) dinosterol, respectively) in the ARC11-R1 core distinguishing between different spring/summer sea ice condition zones. The gradient from dark to light red represents periods of 1820-1930, 1930-1980, and 1980-2020, respectively.
After 1930s, \( P_{III} \) gradually decrease to ca 0.7 while \( P_B \) and \( P_D \) drop to lower values around 0.4 till the 1960s to then slightly increased until the 80s -90s when \( P_B \) and \( P_D \) exceed the seasonal sea ice threshold value (0.5) and \( P_{III} \) that of nearly permanent sea ice at approximately 0.75. In this time interval (1930s - 1980s), low \( IP_{25} \) and increasing brassicasterol in cores ARC4-C07 and ARA01B-03MUC suggest enhanced sea ice melting and the northward retreat of the summer ice edge (Fig. S5). From 1980s to present, \( P_B \) and \( P_D \) continued to decrease but at a faster rate (Fig. 5 and 7a, b, c) emphasizing the unprecedented decline of seasonal sea ice over the last 30 years, as also found by Astakhov et al. (2019) and shown by remote sensing data (Walsh et al., 2017, Wang et al., 2019b).

**Figure 7.** Downcore profiles of (a) \( P_{BIP_{25}} \) index using brassicasterol (\( P_B \)), (b) dinosterol (\( P_D \)) and (c) HBI-III (\( P_{III} \)) in the core ARC04-C07 (green, Bai et al. 2022), ARA01B-03MUC (yellow, Kim et al. 2019) and ARC11-R1 (blue). i, ii and iii in (a), (b) and (c) represent different sea ice cover scenarios, which are schematically illustrated in (d). The color of sea ice in (d) represents the different sea ice cover: light blue for spring/autumn sea ice, and dark blue for summer sea ice.

In summary, the downcore profiles of seasonal sea ice proxies over the last 200 years evidence (Fig. 7d): i) reconstructed lower sea ice cover contrasting with nearly permanent sea ice observation record between the 1820s and 1930s, implying the limitation of \( P_{IP_{25}} \) index under nearly permanent sea ice; ii) from 1930s to 1980s, the seasonal sea ice slowly retreated to the north and the summer sea ice edge gradually reached the location of the R1 core; iii) a strong reduction sea ice cover with summer sea ice.
edge conditions being reached since the 1980s at our core site. Finally, our findings highlight the need to further investigate the potential value of HBI-II in the Arctic Ocean to track drifting and landfast ice from coastal regions under freshwater discharge influence.

325 5.2 Organic carbon variability in response to sea ice change

5.2.1 Modern sources of organic carbon

The loss of sea ice is the most remarkable manifestation of global warming in the Arctic that has profound impacts on the carbon cycle. The most obvious one is the shift of the primary production pattern as a result of light and nutrient supply changes. Enhanced riverine inputs of terrestrial organic matter also affect the nature and amount of organic carbon reaching the Arctic Ocean. In this section, we investigate OC compositional changes over the past 200 years in relation with sea ice conditions by distinguishing three major carbon pools, e.g., sympagic, pelagic and terrestrial. We use the H-print ratio as a complementary indicator to PIP to discriminate pelagic and sympagic marine sources (Brown et al., 2014b; Brown and Belt, 2017; Koch et al., 2020). As shown in figure 8a, high values of H-print (>80%) occur in the western CS where marine phytosterols are high (dinosterol concentrations >5 μg g\(^{-1}\) TOC and brassicasterol concentrations >100 μg g\(^{-1}\) TOC, Su et al., 2022). Intermediate values (30%-60%) are roughly lying between the 1979 and 2019 isolines of the September minimum ice edge, denoting mixed pelagic and sympagic productions (Fig 8a). Minimum H-print values are found in coastal sediments along the western ESS and suggest freshwater discharge, by lowering water salinity, can suppress HBI-II production (Su et al., 2022) (Fig. 8a). At higher latitudes, low H-print values are likely reflecting light limitation due to nearly permanent sea ice cover (Fig. 8a).

Earlier studies in Arctic Eurasia estuary sediments have reported \(\delta^{13}C_{org}\) values ranging from -27.8% to -24.7% (Bröder et al., 2019; Tesi et al., 2014). Lowest values (-27.5% to -26%, Fig. 8b) are found in the western ESS shelf sediments receiving land-derived material from the Indigirka and Lena rivers in agreement with previous data (-27.5% to -25.5%, Bröder et al., 2019). Apart from permafrost thawing, sea ice retreat likely accelerated coastal erosion contributing to the transfer of allochthonous material towards the Arctic Ocean (Overeem et al., 2011). The \(\delta^{13}C_{org}\) values of organic matter produced by phytoplankton in the Arctic Ocean vary around -24 ± 3% (Stein et al., 2004; Vonk et al., 2012), which are similar to those found in saline waters of the western CS characterized by high pelagic production.
Enriched $\delta^{13}C_{org}$ values (-22‰ to -20.5‰) are also observed in the marginal ice zone/seasonal sea ice zone of the CS (Fig. 8b) where sea ice plankton production is significant, which is in accordance with values reported in sea ice (-23.6‰ to -18.3‰) (Schubert and Calvert, 2001).

$\delta^{13}C_{org}$ and H-print were combined to discriminate among autochthonous and allochthonous sources of OC in our surface sediments. As illustrated by the scatter plot in figure 8c, a three-end-member distribution emerged from the relationship between these 2 parameters and summer sea ice concentration (SuSIC) demonstrating their potential to differentiate sympagic, pelagic and terrestrial OC in our dataset. The site symbolizing the sympagic end-member is located at the sea ice edge where sea ice phytoplankton production is high. Coincidentally, its $\delta^{13}C_{org}$ value (-20.5‰) falls within the range of values reported in the lowermost 10 cm of sea ice cores (-23.6‰ to -18.3‰, Schubert and Calvert, 2001) and close to the mean $\delta^{13}C_{org}$ value of sedimentary IP$_{25}$ (-19.3‰±2.3‰, Belt et al., 2008). The terrigenous end-member is consistently represented by sites located in estuarine zones or where land-derived supply dominates. Pelagic end-member values (-23.5‰) are encountered in the western CS where phytoplankton productivity is the highest and the influence of terrigenous and sea ice carbon less pronounced. This more depleted value as compared to IP$_{25}$ is consistent with the latter being produced in sea ice. Indeed, the $\delta^{13}C_{org}$ of the two marine components also depends on $pCO_2$. Higher $pCO_2$ in surface waters may result in lower $\delta^{13}C_{org}$ in pelagic marine algae OC while sea ice diatoms are relatively enriched in $\delta^{13}C_{org}$ due to potentially limited $CO_2$ in sea ice (Tortell et al., 2013). It should be pointed out that our surface sediment dataset covers the nearshore, shelf and basin regions, with different deposition rates therefore the OC source apportionment reflect several tens to hundred years, depending on the location (Bröder et al., 2016; Baskaran and Naidu, 1995; Vonk et al., 2012; Li et al., 2020). Nevertheless, H-print, $\delta^{13}C_{org}$ and SuSIC successfully discriminate OC sources.

Based on these results, a ternary mixing model was used to calculate the relative contribution of sympagic, pelagic, and terrestrial sources in our surface sediments (Fig. 8c and Fig. S6; Table S1) using the following equation:

$$\delta^{13}C_{sample} = f_{sym} \times \delta^{13}C_{sym} + f_{pela} \times \delta^{13}C_{pela} + f_{terr} \times \delta^{13}C_{terr}$$  \hspace{1cm} (4)

$$H - print_{sample} = f_{sym} \times H - print_{sym} + f_{pela} \times H - print_{pela}$$  \hspace{1cm} (5)

$$f_{sym} + f_{pela} + f_{terr} = 100\%$$  \hspace{1cm} (6)
where \( f_{\text{sym}}, f_{\text{pela}}, \) and \( f_{\text{terr}} \) are the sympagic, pelagic, and terrestrial fractions of OC, respectively. The \( \delta^{13}\text{C}_{\text{org}} \) end-members for used for the sympagic end-member at 0% for H-print is -20.5‰. The pelagic \( \delta^{13}\text{C}_{\text{org}} \) end-member at 100% for H-print is -23.5‰. The terrestrial \( \delta^{13}\text{C}_{\text{org}} \) end-member is -27.5‰.

5.2.2 Source and burial of organic carbon change over the last two centuries

In this section, we examine the temporal evolution of sympagic, pelagic and terrestrial fractions of OC calculated with our ternary mixing model and end-member values, with changing sea ice conditions over the last two centuries. Figure 9a shows that TOC values slowly increase between 1820 and 1860 and remain relatively stable around 0.8% until 1960 where they increase rapidly to reach their highest levels (1.17%) at the core-top, falling in the range reported in the surface sediments of the CS shelf and CP (1.25% to 2.56%, Goñi et al., 2013; 0.31% to 1.73%, Ji et al., 2019). This trend is paralleled by increasing \( \delta^{13}\text{C}_{\text{org}} \) values (Fig. 9a), suggesting that higher TOC may be related to enhanced marine production (primary and secondary). Indeed, sea ice retreat and increase ice free conditions are expected to result in higher rates of primary production due to higher light penetration and more nutrients supply both from river and via wind driven mixing. In addition, a longer season of production will also subsequently enhanced production and export to the sea floor (Ouyang et al., 2022; Zhuang et al., 2022).
Figure 8. Distribution of (a) H-print and (b) $\delta^{13}$C$_{org}$ in the surface sediments of the ESS and CS. (c) Cross-plot of values of $\delta^{13}$C$_{org}$ and H-print for surface sediments. The gradient from dark blue to white represents SuSIC (Summer Sea Ice Concentration, NSIDC) from 0-20%, 20-40%, 40-60%, 60-80% and 80-100%, respectively, and ARC11-R1 core (in orange). The $\delta^{13}$C$_{org}$ end-members for the sympagic end-member at 0% for H-print is -20.5‰. The pelagic $\delta^{13}$C$_{org}$ end-member at 100% for H-print is -23.5‰. The terrestrial $\delta^{13}$C$_{org}$ end-member is -27.5‰. The dotted and dashed lines in black represent the 20% isolines of September sea ice concentration for the 2019 and 1979, respectively.
Table 3. Overall organic carbon composition of core ARC11-R1 (relative portion, \( f_{\text{occ}} \) (%), and absolute content, OC (mg g\(^{-1}\) d. w.)).

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<th>( f_{\text{terr}} ) (%)</th>
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The C/N ratios are higher than the Redfield ratio over the entire core and show a decreasing trend (Fig. 9b), in agreement with diminishing contribution of terrestrial with respect to marine OC sources. The mean \( f_{\text{terr}} \) is relatively high throughout the R1 core (46.45% ± 8.20%, Fig. 9c and Table 3) but decrease from 58.70% to 30.45%. Although R1 is located rather far from the continent, these estimates suggest efficient transport pathways of land-derived organic matter towards the open sea such as drifting sea ice (Jia et al., 2020) or wave action remobilizing and transporting shelf sediments to the ocean interior (Vonk et al., 2012). Time series sediment traps evidenced high and long-standing terrigenous material advection by lateral transport controlled by Chukchi Slope Current and mesoscale eddies in winter in this region (Onodera et al., 2021; Watanabe et al., 2014; Watanabe et al., 2022). Figure 9c also shows that since the 1960s, there has been a significant decline in the share of terrigenous OC with the enhanced loss of sea ice of the last 60 years, and subsequent rise of productivity of marine and sea ice diatoms. By contrast, ARC4-C07, the core located to the southeast of R1 (Fig. 1), exhibits a trend of enhanced land-sourced input in recent years (Bai et al., 2022). The different trend of the terrigenous OC components of these two cores might be related to the weakening of sediment-laden sea ice transport to higher northern latitudes due to its earlier melting.
From the 1820s to 1930s, pelagic and sympagic OC are found in comparable amounts in the sediments (2.17 ± 0.53 mg g⁻¹ d. w. sediment and 1.72 ± 0.46 mg g⁻¹ d. w. sediment, respectively; Fig. 9d and Table 3), but from the 1930s to 1980s, the content of sympagic OC increase by nearly 50% to 3.10 mg g⁻¹ d. w. sediment (Fig. 9d). However, pelagic phytoplankton growth also shows rising values but remain lower than sympagic OC possibly because of surface freshening limiting nutrient supply (Arrigo et al., 2008). After the 1980s, when the minimum ice edge reached R1 and the ice-free period prolonged in the northern CS (Astakhov et al., 2019), the sympagic and pelagic productions depict a final increase and superseding terrestrial OC (Fig. 9c and 9d). The high δ¹⁵N data during this period suggest enhanced stratification most likely due to surface freshening (Fig. 3d) which gradually shifted the limiting factor of marine pelagic production from light to nutrients availability (Ardyna and Arrigo, 2020;
6 Conclusions

H-print and δ^{13}C_{org} values from 83 surface sediments were used to diagnose OC sources across the western Arctic Ocean and how they are influenced by sea ice extent. δ^{13}C_{org} values were consistently generally lower in the southern coastal and shelf sediments where riverine terrigenous inputs are stronger, while heavier δ^{13}C_{org} were found at offshore sites of the northern CS and ESS where pelagic and/or sympagic productions are significant. Combining δ^{13}C_{org} and H-print enabled to draw an inventory of terrestrial, sympagic and marine pelagic OC in northern CS sediments using a ternary mixing model which was then applied to the ARC11-R1 core to reconstruct the temporal evolution of the OC content since 1820 as the sea ice retreat.

Our results demonstrate that over the last 200 years the northern CS experienced nearly permanent sea ice conditions between 1820s and 1930s followed by a period of gradual melting to reach marginal sea ice conditions (1930s-1980s). The most recent decades were marked by the accelerated decline of sea ice (1980s - present) leading to shifts in primary production that, in turn, impacted the composition and burial of OC in Arctic sediments from coastal to open sea areas. Our data show that with the loss of sea ice, the fraction of terrigenous OC decreased while marine pelagic and sympagic OC export and sequestration in the deep-ocean increased. Since the beginning of the 21st century the three OC pools at the core site are comparable in size while 200 years ago OC was predominantly of terrestrial origin. In the future, with rising temperature and reduced sea ice, primary production in the northern CS may however become nutrients limited as a result of freshening and ocean stratification, and phytoplankton populations will likely undergo further alteration with subsequent changes in CO₂ drawdown.
Data availability.

All data that support the findings of this study are included within the article and supplement.

Author contributions.

L.S., J.R. and J.C. designed the study and wrote the manuscript with contribution of M.-A.S., Y.B., Z.L., R. Z., H.J., A.A.S. and X.S. L.S. and R. Z. contributed the biomarker analyses and the determination of bulk parameter. J.R. retrieved the environmental data from different database while X.H. carried out the age model estimate of R1. All authors contributed to the final version of the manuscript.

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

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