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Changes in terrestrial organic matter input to the Mendeleev Ridge, western Arctic Ocean, during the Late Quaternary

Masanobu Yamamoto a*, Leonid Polyak b,

a Faculty of Environmental Earth Science, Hokkaido University, Kita-10, Nishi-5, Kita-ku, Sapporo 060-0810, Japan
b Byrd Polar Research Center, Ohio State University, 1090 Carmack Road, Columbus, OH 43210, USA
* Corresponding author. E-mail address: myama@ees.hokudai.ac.jp (M. Yamamoto).

Abstract

Hydrocarbons and glycerol dialkyl glycerol tetraethers (GDGTs) were analyzed in Late Pleistocene sediment s of Core HLY0503-08JPC collected at the Mendeleev Ridge during the Healy-Oden Trans Arctic Expedition 2005 (HOTRAX’05) to investigate environmental changes in the western Arctic Ocean during the last full glacial cycle, ca. 130 kyr. Variations in long-chain n-alkane and GDGT concentrations correspond to alternated color banding, brown (interglacial/interstadial) and grayish (glacial/stadial) layers. Grayish layers are characterized by abundant higher plant n-alkanes and branched GDGTs, implying larger contribution of terrestrial plant and soil organic matter (OM) in glacial environments, possibly due to the deposition of fine-grained products of glacial erosion in the Amerasian basin. Lithic n-alkanes derived from mature OM show pronounced peaks, which can be classified into six types presumably indicative of various sediment sources. Some peaks are correlated to events of iceberg discharge and freshwater outbursts from proglacial lakes of the Eurasian and, possibly, Laurentide ice sheets, suggesting that other peaks may correspond to similar events.

Keywords; Arctic, biomarker, tetraethers, glacial, HOTRAX’05, ice rafted debris, paleoenvironment.

1. Introduction

The Arctic Ocean plays a major role in global climate changes by changing global energy balance through ice-albedo feedback and by affecting the oceanic thermohaline circulation through the water exchange with the Atlantic and Pacific oceans (Aagaard and Carmack, 1989; Delworth et al., 1997; Smith et al., 2002). A complete understanding of the role of the Arctic Ocean in global climate changes requires records of the formation and coverage of iceberg and sea ice during past periods.

Icebergs and sea ice deliver ice-rafted debris (IRD), which has been used to reconstruct the Arctic’s iceberg and sea ice formation history (e.g., Bischof et al., 1996; Polyak et al., 2004; Spielhagen et al., 2004). IRD source areas have been determined using mineral and chemical compositions (e.g., Stein et al., 1994;
Bischof and Darby, 1997; Darby et al., 2002). Organic geochemistry has also been applied in such studies; the amount and quality of bulk OM have been reported for sediment cores from the Amundsen Basin (Schubert and Stein, 1996) and the Siberian continental margins (Stein et al., 2001, 2004; Bouscein et al., 2002). Biomarkers are also used as source-specific tracers in oceanographic studies in the central Arctic Ocean (Schubert and Stein, 1997; Belicka et al., 2002; Yunker et al., 2005), the northern Eurasian margin (Peulvé et al., 1996; Fahl and Stein, 1997; Zegouagh et al., 1996) and North American margins (Yunker et al., 1995; Belicka et al., 2004), and in paleoceanographic studies in Holocene and late Pleistocene sequences of the Laptev and Kara Seas (Fahl and Stein, 1999; Stein et al., 2001), and the Lomonosov Ridge in the central Arctic (Yamamoto et al., 2008).

Detrital matter is mostly transported to the central Arctic Ocean by ice rafts; fine sediment can be also carried by intermediate-depth currents. Under present conditions the Transpolar Drift transports sediment-laden “dirty” sea ice from the Laptev Sea; eastward boundary currents along the northwestern Siberian margin and clockwise water circulation in the Beaufort gyre join with the Transpolar Drift (Gordienko and Laktionov, 1969; see Fig. 1) to contribute to mass transportation by ice-drift streams (Nürnberg et al., 1994).

We examined late Pleistocene biomarker records collected from the 2005 Healy-Oden Trans Arctic Expedition (HOTRAX’05) at the Mendeleev Ridge in the western Arctic Ocean, with the goal of understanding glacial-interglacial variation in biomarker composition in the Arctic Ocean and their relationship to late Pleistocene climate change. The results indicate drastic changes in biomarker composition with glacial–interglacial variation, which is discussed in relation to changes in sediment delivery mechanisms and source areas, and the expansion and disintegration of the Eurasian and North American ice sheets.

2. Materials and methods

2.1. Samples

The Mendeleev Ridge is an elevated portion of the Arctic Ocean floor extending northwards from the East Siberian shelf. Core HLY0503-08JPC, ca. 12 m long, was collected on the Mendeleev Ridge (79°35.6’N, 172°30.1’W) at a water depth of 2792 m in the western Arctic Ocean on the 2005 Healy-Oden Trans Arctic Expedition (HOTRAX’05) (Darby et al., 2005; Fig. 1).

The core site is located close to the area of interaction of the two major Arctic Ocean circulation systems, the Beaufort Gyre and the Transpolar Drift (Fig. 1), and is thus affected by deposition of both North American and Eurasian sediment rafted by sea ice and/or icebergs (Adler et al., this volume). In addition, fine-grained material may be transported by water current that flows at intermediate water depths along the Eurasian slope. Comparison with other cores from the Mendeleev Ridge collected during HOTRAX’05 and earlier expeditions (Clark et al., 1980; Darby et al., 1997; Poore et al., 1999; Polyak et al., 2004) indicates that Core HLY0503-08JPC has relatively elevated sedimentation rates resulting probably from its location at a local depocenter at the bottom of the eastern flank of the ridge (Adler et al., this volume). Enhanced sedimentation rates combined with a strategic location at a junction of two
major circulation and sedimentation pathways makes Core HLY0503-08JPC well suited for a detailed investigation of sedimentary environments in the Amerasian Basin of the Arctic Ocean.

Sediments comprise mud and sandy mud with strong color banding mainly “grayish” layers (yellowish brown to olive brown) and “brown” layers (brown to dark brown) (Adler et al., this volume). Color bands range in thickness from 2 to 35 cm and typically have bioturbated bottom contacts. In addition to this cyclic lithostratigraphy, there are thin intervals of pinkish and/or whitish coloration due to enrichment of sediment with carbonate clasts, especially conspicuous at ~0.45 m, ~1.15 m and ~2.3 m. This pattern of sequence can be recognized throughout the entire western Arctic Ocean (Phillips and Grantz, 2001; Polyak et al., 2004). Various paleoceanographic proxies measured in this core such as the diffuse spectral reflectance in the visual spectra (exemplified by L* and a* channels), the content of sand and coarser fraction (>63 μm), and the δ18O and δ13C in the calcite of planktonic foraminifer Neogloboquadrina pachyderma (sinistral) are reported in Adler et al. (this volume).

Cores were stored at ~5°C. Samples were taken every 1 cm from the uppermost 250 cm of the core and immediately frozen at -20°C. Samples were freeze-dried, mixed and split into aliquots. A total of 250 samples was analyzed in this study.

2.2. Age–depth model

We used the age-depth model generated by Kaufman et al. (2008) and Adler et al. (this volume). The age-depth model was developed from ten C-14 dates of planktonic foraminifer Neogloboquadrina pachyderma (sin.) (4.0-44.8 ka), twenty-three dates by the amino-acid racemization rates in N. pachyderma (sin.), six tie points obtained from stratigraphic correlation with ACEX Holes and adjacent cores from the central Lomonosov Ridge (O’Regan et al., 2008), and five tie points obtained from correlation of glacial units with Eurasian glaciations (Svendsen et al., 2004; Larsen et al., 2006). According to this age model, the sequence of the uppermost 250 cm covers the last ca. 130 ka (MIS-1 to MIS-5e).

2.3. TOC measurement

Samples were analyzed for total organic carbon (TOC) using a LECO WR-112 carbon analyzer. The analyzer was attached to a halogen trap (antimony and potassium iodide). To remove carbonate carbon, the sample was acidified as follows: aliquots of ca. 0.5–1 g were soaked in 1 M HCl in a ceramic crucible overnight and then heated at 150°C for 3 h after adding more 1 M HCl. The sample was rinsed (x 2) to remove chlorides by adding distilled water and was again heated at 110°C for 3 h. The measurement precision was better than 0.01 wt%.

2.4 Hydrocarbon analysis

Lipids were extracted (x 3) from ca. 5 g of dried sediment using a DIONEX Accelerated Solvent Extractor ASE-200 at 100°C and 1000 psi for 10 min with 11 ml of CH2Cl2–CH3OH (6:4) and then concentrated. The lipid
extract was separated into four fractions using column chromatography (SiO$_2$
with 5% distilled water; i.d., 5.5 mm; length, 45 mm): F1 (hydrocarbons), 3 ml
hexane; F2 (aromatic hydrocarbons), 3 ml hexane-toluene (3:1); F3 (ketones), 4
ml toluene; F4 (polar compounds), 3 ml toluene–CH$_3$OH (3:1); $n$-C$_{36}$H$_{74}$ and
C$_{46}$ GDGT (2,2'-di-0-decyl-3,3'-di-O-(1',ω'-eicosanyl)-1-1'-di-(rac-glycerol);
Patwardhan and Thompson, 1999) were added as internal standards to F1 and F4,
respectively.

Gas chromatography (GC) was conducted using a Hewlett Packard 6890
gas chromatograph with on-column injection and a flame ionization detector
(FID). Samples were dissolved in hexane. He was the carrier gas and the flow
velocity was maintained at 30 cm/s. A Chrompack CP-Sil5CB column was used
(length, 60 m; i.d., 0.25 mm; thickness, 0.25 μm). The oven temperature was
programmed to rise from 70 to 130°C at 20°C/min, from 130 to 310°C at
4°C/min, and to hold at 310°C for 30 min. The standard deviations of five
duplicate analyses averaged 7.5% of the concentration for each compound.

2.5. Glycerol dialkyl glycerol tetraether (GDGT) analysis

An aliquot of F4 was dissolved in hexane–2-propanol (99:1). Glycerol
dialkyl glycerol tetraethers (GDGTs) were analyzed using high performance
liquid chromatography-mass spectrometry (HPLC-MS) with an Agilent 1100
HPLC system connected to a Bruker Daltonics micrOTOF-HS time-of-flight
mass spectrometer. Separation was conducted using a Prevail Cyano column
(2.1 x 150 mm, 3μm; Alltech) and maintained at 30°C following the method of
Hopmans et al. (2000) and Schouten et al. (2007). Conditions were: flow rate
0.2 ml/min, isocratic with 99% hexane and 1% 2-propanol for the first 5 min
followed by a linear gradient to 1.8% 2-propanol over 45 min. Detection was
achieved using atmospheric pressure, positive ion chemical ionization-mass
spectrometry (APCI-MS). The spectrometer was run in full scan mode ($m/z$
500–1500). Compounds were identified by comparing mass spectra and
retention times with those of GDGT standards (formed from the main
phospholipids of Thermoplasma acidophilum via acid hydrolysis) and those in
the literature (Hopmans et al., 2000). Quantification was achieved by integrating
the summed peak areas in the (M+H)$^+$ and the isotopic (M+H+1)$^+$ ion traces and
comparing these to the peak area of an internal standard (C$_{46}$ GDGT;
Patwardhan and Thompson, 1999) in the (M+H)$^+$ ion trace, according to the
method of Huguet et al. (2006). The correction value of ionization efficiency
between GDGTs and the internal standard was obtained by comparing the peak
areas of T. acidophilum-derived mixed GDGTs and C$_{46}$ GDGT of known
amounts. The standard deviation of a replicate analysis is 3.0% of the
concentration for each compound.

3. Results and discussion

Typical gas chromatograms of F1 and HPLC-MS chromatograms of F-4
are shown in Figs. 2 and 3, respectively.

3.1. TOC content

The TOC content varies between 0.02 and 0.48%, with an average of
0.28%. The TOC decreases exponentially with increasing depth (Fig. 4D).

3.2. Normal-alkane distribution

N-alkanes occur as a major component of the F1 (hydrocarbon) fraction and show a unimodal distribution with a maximum at C\textsubscript{25}-C\textsubscript{31} (Fig. 2). The odd carbon number preference index (CPI) values of C\textsubscript{24}-C\textsubscript{34} homologues (Bray and Evans, 1961) vary between 1.5 and 5.3, with an average of 4.0 (Fig. 4F). The average chain length (ACL) values of C\textsubscript{27}-C\textsubscript{33} homologues (Bray and Evans, 1961) vary between 29.2 and 29.8, with an average of 29.6 (Fig. 4F). Most samples showed n-alkane distribution of mixture of a pattern with high odd carbon number preference (CPI = ~5; Fig. 2A) and high average chain length (ACL = ~29.7) and a pattern heavily influenced by mature (thermally altered) n-alkane with low CPI (~1.5; Fig. 2B) and low ACL (~29.2). The former is typical of terrestrial higher plant waxes (Eglinton and Hamilton, 1967). This pattern indicates that the OM was derived from fresh higher plant material. The latter is typical of coal and coaly shale (Bray and Evans, 1961), indicating an influx of mature (lithic) OM.

If we assess the end-member CPI values of higher-plant and mature OM, we can estimate the concentrations of n-alkanes of both origins. Riverine suspended matter and surficial sediments from the Arctic Ob and Yenisei Rivers have CPI values from 3.2 to 8.1 (average 5.8; Fernandes and Sicre, 2000). Assuming that the long-chain n-alkanes derived from higher plant waxes have CPI of 5.2 (maximal value for samples in the study core), and those derived from mature rocks have CPI of 1 (typical values of OM in catagenesis and metagenesis stages; Tissot and Welte, 1984), we calculated concentrations of “higher plant” and “lithic” C\textsubscript{25}-C\textsubscript{33} n-alkanes from total C\textsubscript{25}-C\textsubscript{33} n-alkane concentration and the CPI value using the following formulae:

\[
\text{Higher plant } \text{C}_{25-33} \text{ n-alkane} = \text{total C}_{25-33} \text{ n-alkane} \times \frac{6.2 (\text{CPI} - 1)}{4.2 (\text{CPI} + 1)}
\]

\[
\text{Lithic } \text{C}_{25-33} \text{ n-alkane} = \text{total C}_{25-33} \text{ n-alkane} - \text{higher plant C}_{25-33} \text{ n-alkane}
\]

“Higher plant” n-alkane concentration varies with sediment color approximated by L* (lightness); Higher concentrations are found in high L* intervals (grayish layers), which also have lighter oxygen isotopic composition of planktonic foraminiferal calcite (Figs. 4B and 4E). In contrast, “lithic” n-alkane concentration is not synchronized with L* (Figs. 4B and 4E).

3.2. GDGT distribution

HPLC-MS revealed that GDGTs are a major high-molecular-weight component of the F4 and comprise both isoprenoid and branched components. The isoprenoid GDGTs include cardarchaeol, GDGT-1, GDGT-2, crenarchaeol, and crenarchaeol regioisomer (Fig. 3). Caldarchaeol and crenarchaeol are relatively abundant in isoprenoid GDGTs. Branched GDGTs make up a homologous series with four to six methyl groups (Hopmans et al., 2004; Fig. 3).
Branched GDGTs are presumably derived from anaerobic bacteria and occur widely in soils and peat bogs (Weijers et al., 2006a). Isoprenoid GDGTs are not source specific because they derive from both marine and terrestrial Archaea (Nishihara et al., 1997; Sinninghe Damsté, 2002b). Crenarchaeol in marine sediments was thought to be derived solely from pelagic crenarchaeota (Sinninghe Damsté et al., 2002b). Microbiological studies have demonstrated the presence of marine crenarchaeota in Arctic waters (Bano et al., 2004). Weijers et al. (2006b) recently found crenarchaeol in soil samples. The crenarchaeol found in our samples is associated with abundant branched GDGTs, suggesting possible contribution of soil crenarchaeota. It is thus likely that crenarchaeol in the study samples has a mixed origin of soil crenarchaeota and aquatic crenarchaeota.

Concentrations of both branched and isoprenoid GDGTs exponentially decrease with increasing depth (Fig. 4G). This decreasing trend is attributed to degradation of GDGTs during early diagenesis. Superimposed on this general decreasing trend, the concentrations show a significant fluctuation that corresponds to L* variation (Fig. 4G). The concentration maxima occur in higher L* intervals (grayish layers).

Hopmans et al. (2004) defined the Branched and Isoprenoid Tetraether (BIT) index as the ratio of branched GDGTs to the sum of branched GDGTs and crenarchaeol as an index of soil OM contribution. BIT ranges from ~0.46 to 0.93 with the average of 0.73 (Fig. 4H). The range of values is similar to that in the central Arctic Ocean (~0.5-1; Yamamoto et al., 2008), but is anomalously high for normal pelagic marine sediments. The high BIT index in the western Arctic Ocean reflects a significant contribution of terrestrial soil OM. Yamamoto et al. (2008) reported that major biomarkers in central Arctic sediments (Site M4, Fig. 1) are long-chain n-alkanes, n-fatty acids and n-alkan-1-ols, indicating terrestrial OM input. The dominance of terrestrial biomarkers was attributed to severe OM degradation caused by slow sedimentation in oxygen-rich benthic water and/or low primary production due to permanent sea ice coverage (Yamamoto et al., 2008). The dominance of long-chain n-alkanes and high BIT in Core HLY0503-08JPC is consistent with these earlier observations in the central Arctic Ocean, suggesting that the environment of the study site has been under similar conditions. Low BIT corresponds to low L* (brown layers), indicating higher contribution of crenarchaeol that derived solely from crenarchaeota, which might have affected by the contribution of marine crenarchaeota living in shelf and/or pelagic water.

3.3 Degradation of OM during early diagenesis
Bottom sediments in the central Arctic Ocean are well oxygenated, as demonstrated on the ACEX core from the Lomonosov Ridge (Backman et al., 2006). The pore water chemistry indicates that manganese oxide reduction occurs between 1.5 and 80 m below sea floor, iron oxide reduction occurs between 20 and 200 m below sea floor, and sulfate reduction starts below 200 m below sea floor (Backman et al., 2006). We infer that similar oxidative environments characterize Core HLY0503-08JPC, which is stratigraphically and sedimentologically comparable to cores from the Lomonosov Ridge (Adler et al.,
The oxygenated sediments are ideal for evaluating the degradation rates of organic compounds with time. The general decreasing trends of TOC, C_{25-33} n-alkanes, isoprenoid GDGTs, and branched GDGTs with increasing depth and age are attributed to the degradation of OM during early diagenesis (Fig. 5). We calculated the first-order decomposition rate constants ($k$ [yr$^{-1}$]) and initial concentrations ($C_0$) by fitting concentrations ($C_t$) versus sample ages ($t$ [yr$^{-1}$]) to the following equation:

$$C_t = C_0e^{-kt}$$

The calculated values of first-order decomposition rate constant ($k$ [yr$^{-1}$]) and initial concentration ($C_0$) are shown in Table 1. The $k$ values of branched GDGTs ($2.0 \times 10^{-5}$ yr$^{-1}$) and isoprenoid GDGTs ($2.5 \times 10^{-5}$ yr$^{-1}$) are higher than those of TOC ($1.1 \times 10^{-5}$ yr$^{-1}$) and C_{25-33} n-alkanes ($1.0 \times 10^{-5}$ yr$^{-1}$). Long-chain n-alkanes are as stable as TOC, being consistent with previous observations (Prahl et al., 1997; Sinninghe Damsté et al., 2002a). GDGTs are relatively less stable than TOC and long-chain n-alkanes in oxygenated conditions.

3.4 Changes in the source of OM in glacial and interglacial regimes

Sediments of Core HLY0503-08JPC are characterized by a distinct alteration of brown and yellowish-gray layers interpreted as interglacial/interstadial and glacial/stadial environments, respectively (e.g., Jakobsson et al., 2000; Polyak et al., 2004; Adler et al., this volume). This study shows that grayish layers are characterized by abundant higher plant n-alkanes and branched GDGTs (Figs. 4E and 4G), suggesting that the OM derived from higher plants and soil is more abundant in these layers than brown layers. The concentrations of higher plant n-alkanes and branched GDGTs are the functions of (1) preservation degree of the compounds, (2) climates of source region, (3) transportation mechanisms, and (4) transportation pathway.

Brown layers are thought to be deposited under the conditions of well ventilated intermediate and bottom waters (Jakobsson et al., 2000; Polyak et al., 2004). The resulting oxygenated condition generally enhances the degradation of OM, thus reducing the concentration of organic compounds. In the study core, variation in TOC did not correspond to brown/gray cycles. Because the first-order decomposition rate constant of TOC ($1.1 \times 10^{-5}$ yr$^{-1}$) is the same order of that of C_{25-33} n-alkanes ($1.0 \times 10^{-5}$ yr$^{-1}$; Table 1), little correspondence of TOC to brown/gray cycles implies that changes in redox conditions are not the principal factor regulating the concentrations of C_{25-33} n-alkanes.

The OM derived from higher plants and soil is more abundant in grayish layers deposited in glacial/stadials. The production of higher plants is generally higher in warmer and wetter climates (Whittakker et al., 1970). The concentration of branched GDGTs in soil has positive correlation with mean annual temperature and also has negative correlation with the soil pH (Weijers et al., 2006a), implying that the production of branched GDGTs are higher in warmer and wetter climates. However, higher concentration of branched GDGTs in the Arctic cores occurs in glacial/stadial rather than interglacial/interstadial intervals. Thus, the larger contribution of higher plant and soil OM in the study
samples does not reflect climates of source regions, but more efficient transportation of OM from land to the Arctic Ocean.

Concentrations of higher plant n-alkanes and branched GDGTs in grayish layers are not related with the content of coarse fraction (>63 µm; Figs. 4C, 4E, and 4G). This suggests that higher plant and soil OM were transported not with coarse particles, but rather with fine particles either by sea ice rafting or subsurface water currents. Fine, muddy sediment in the central part of grayish layers suggests negligible contribution from ice rafting, probably due to the solidly ice-bound ocean surface (Polyak et al., 2004; Adler et al., this volume). Polyak et al. (this volume) suggest that the major source of this sediment was glacial flour delivered by meltwater from glaciated margins at the Arctic Ocean periphery. Abundance of higher plant n-alkanes and branched GDGTs in grayish layers is thus related to the deposition of fine-grained products of glacial erosion. The peat that was deposited in northern North America and Eurasia during interglacials is a potential source of higher plant and soil OM. Ice erosion of peat during glacial periods could deliver such an OM into the Amerasian Basin. Because this hypothesis is still speculative, we need further provenance studies to clarify the source and transportation mechanisms of OM.

3.5 Intermittent transport of mature OM by icebergs and sea ices

Concentration of lithic n-alkanes and the relative abundance of lithic n-alkanes in total n-alkanes (reflected as CPI reversely) show sixteen maxima in the uppermost 250 cm of the study core (Fig. 6). Ten of sixteen peaks occur at the boundary between grayish and brown layers (Fig. 6). Eight of sixteen peaks have maximal peaks of the content of coarse fraction (> 63 µm). Another series of eleven peaks have minimal peaks of ACL. Based on associated peaks of coarse fraction (> 63 µm), detrital carbonate layers and ACL, the maximal peaks of lithic n-alkanes are classified into six different types (Types A-F in Fig. 6).

The peaks of type A are associated with maximal sand content and occur at or close to detrital carbonate layers. Because the only source of this detrital carbonate (mostly dolomite) in the central Arctic Ocean is derived from the Laurentide part of the Canadian Archipelago (Bischof et al., 1996; Darby et al., 2002), this type may have derived by icebergs from the same source.

The peaks of type B are associated with minimal ACL and high content of coarse fraction (> 63 µm). The peak at 247.5 cm in depth belonging to this type correlates to a major peak of iron oxide grains derived from the Innuitian ice sheet in NP26 cores (Polyak et al., 2004). This correspondence suggests that this type might have derived by icebergs from the Innuitian ice sheet.

The peak of type C is associated with a strong minimum in ACL and a maximal content of coarse fraction (> 63 µm) and correlates to a major peak of quartz IRD and iron oxide grains attributed to Barents and Kara sources in NP26 cores (Polyak et al., 2004). The stratigraphic position of this peak corresponds to a large glaciation in northern Eurasia associated with freshwater discharge from large ice-dammed lakes at ~75 ka (Spielhagen et al., 2004).

The peaks of type D are associated with strong minima in ACL. The peak at 215.5 cm in depth belonging to this type correlates to a major peak of iron oxide grains attributed to the Laptev and East Siberian seas in NP26 cores.
(Polyak et al., 2004), suggesting northern Siberian origin.

The peaks of type E are associated with moderate minima in ACL. The peak at 66.5 cm in depth is associated with a $\delta^{18}O$ depletion of planktonic foraminifer (Figs 4B and 4F), which is attributed to a voluminous pulsed meltwater discharge (Adler et al., this volume). The peaks at 128.5 and 141.5 cm in depth correlate to major peaks of iron oxide grains attributed to the Barents and Kara seas in NP26 cores, and the peak at 172.5 cm in depth correlates to that of the Laptev and East Siberian seas (Polyak et al., 2004), suggesting northern Siberian origin.

The peak of type F is associated with a non-carbonaceous IRD spike. The stratigraphic position of this peak corresponds to a large glaciation in northern Eurasia associated with freshwater discharge from large ice-dammed lakes at ~52 ka (Spielhagen et al., 2004).

Preliminary correlation between the types of lithic OM peaks in the study core and the types of iron oxide grains in NP26 cores suggests that the type of lithic OM is also useful for identifying the sediment source. Future studies such as a more detailed finger-print analysis for trace biomarkers and a comparison with the composition of iron oxide grains have a potential for refining the provenance of the Arctic Ocean sediments.

Some of minimal peaks of CPI at the boundary between gray and brown layers show a saw-tooth changing pattern of CPI; the sudden drops of CPI and the subsequent gradual increases (Fig. 6). This implies that mature OM was massively delivered at the initial stage of each event and was gradually replaced by higher plant and soil OM. Spielhagen et al. (2004, 2005) and Jakobsson et al. (2007) suggested that the outburst of the water from ice dammed lakes in Siberia occurred several times during the Late Pleistocene. Similar events may have taken place on the North American side, but they are less understood except for the last deglaciation (Poore et al., 1999; Polyak et al., 2007). Such an outburst combined with a massive iceberg discharge could have eroded bedrocks including coals and coaly shales and discharged mature OM onto the continental shelf. After that, gradual supply of higher plant and soil OM by rivers could dilute mature OM preserved on the shelf. This mechanism could change the composition of OM in the source area from lithic to higher plant and soil OM.

5. Conclusions

Biomarker records from western Arctic Ocean sediments show drastic changes with glacial–interglacial variation. Concentrations of higher plant-derived compounds, thermally altered compounds, and soil-derived compounds are highly sensitive to past changes in sediment delivery mechanisms and source areas.

Variations in long-chain n-alkanes and GDGT concentrations indicate higher contribution of terrestrial OM (higher plant and soil OM) during glacial/stadials. These changes in the source of OM are presumably related to the enhanced delivery of the products of glacial erosion including higher plant and soil OM from the glaciated margins into the Amerasian basin.

Concentration maxima of lithic n-alkanes derived from mature OM may correspond to events of iceberg discharge and freshwater outburst from
proglacial lakes that carried eroded bedrock material.

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Table 1. First-order decomposition rate constants ($k$) and initial concentrations ($C_0$) of TOC, long-chain n-alkanes, and isoprenoid and branched GDGTs

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<td>TOC</td>
<td>0.34%</td>
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<td>$C_{25-33}$ n-alkanes</td>
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<td>1.0</td>
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<td>Isoprenoid GDGTs</td>
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<td>Branched GDGTs</td>
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Figure captions

Fig. 1. Locations of study site (8JPC; Darby et al., 2005) and ACEX site (M4; Backman et al., 2006) and the bathymetry and surface currents of the Arctic Ocean.

Fig. 2. Gas chromatograms of F1 in samples at (A) 1-2 cm (B) 141-142 cm. Numbers indicate carbon numbers of n-alkanes. IS is an internal standard (n-C_{36}H_{74}).

Fig. 3. The HPLC-MS base peak of tetraether lipids in the F4 fraction of sample at 1-2 cm.

Fig. 4. Depth variations in (A) age (Kaufman et al., 2008; Adler et al., this volume), (B) color reflectance L* and $\delta^{18}$O of 150-250 $\mu$m planktonic foraminifera *N. pachyderma* (sin.) (Adler et al., this volume), (C) content of sand and coarser fraction (> 63 $\mu$m particles; Adler et al., this volume), (D) total organic carbon (TOC) content, (E) concentrations of total, and higher plant long-chain C_{25}-C_{33} n-alkanes, (F) carbon number preference index (CPI) and average chain length (ACL) of C_{27}-C_{33} n-alkanes, (G) concentrations of branched and isoprenoid GDGTs, and (H) the branched and isoprenoid tetraether (BIT) and the ratio of caldarchaeol to crenarchaeol (Cald/Cren) in Core FLY0503-08JPC. Shades indicate brown sediment layers; dark shades show the most conspicuous detrital carbonate layers. Brown and grayish units are indexed as in Polyak et al. (2004).

Fig. 5. Age variations in TOC content, long-chain C_{25}-C_{33} n-alkane concentration, and concentrations of branched and isoprenoid GDGTs in Core HLY0503-08JPC.

Fig. 6. Depth variations in color reflectance L* and the content of sand and coarser fraction (> 63 $\mu$m particles) (Adler et al., this volume), lithic long-chain C_{25}-C_{33} n-alkane concentration, and CPI and ACL of C_{27}-C_{33} n-alkanes in Core HLY0503-08JPC. Letters of A-F indicate the types of peak. Single star and double stars indicate moderate and strong peaks, respectively.
Fig. 1
Fig. 2
Fig. 4
Fig. 5
Fig. 6