Data report: no alkenones detected by shore-based GC-MS analyses of Eocene samples from Site U1356

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Abstract

We applied shore-based gas chromatography–mass spectrometry (GC-MS) to confirm shipboard analyses of samples from Integrated Ocean Drilling Program (IODP) Expedition 318. Initially, 25 samples from Site U1356, ranging in age from early Eocene to late Miocene, were extracted for lipids, separated into chemical fractions, and analyzed by GC-MS onboard the R/V JOIDES Resolution. Molecular biomarker classes identified include aliphatic hydrocarbons, hopanes, unresolved complex mixtures of branched alkanes, ketones, n-alkanols, sterols, diols, and n-alkanoic acid methyl esters. Relative to the Oligocene and Miocene, molecular concentrations generally increase by several orders of magnitude in the early Eocene sediments. Additional shore-based analyses confirmed the presence of a diverse range of terrestrial biomarkers that includes plant-waxes and hopanes. However, additional analysis of the ketone fraction from 50 Eocene samples from Site U1356 detected no alkenone compounds. This negative result supersedes the initial report of alkenones in the Eocene sediments from Site U1356 and indicates that shipboard assignment of alkenones was erroneous.

Introduction

Integrated Ocean Drilling Program (IODP) Expedition 318 to the Wilkes Land margin of Antarctica recovered a sedimentary succession ranging in age from the early Eocene to the Holocene. Site U1356 is at the transition between the continental rise and the abyssal plain at 4003 meters below sea level (mbsl) and was cored to 1006.4 meters below seafloor (mbsf) (see Figure F16 in the “Site U1356” chapter [Expedition 318 Scientists, 2011b]). The sediments are divided into 11 lithostratigraphic units, and age control is based on biostratigraphic datums from siliceous microfossils (diatoms and radiolarians), calcareous nannofossils, and organic-walled dinoflagellates (dinocysts) integrated with a magnetostratigraphy that is correlated, where feasible, to the geomagnetic polarity timescale (GPTS) (Tauxe et al., 2012). This data report is concerned with samples from the lowermost 110 m of Site U1356 (lithostratigraphic Units X and XI), dated to the early to middle Eocene (53.6–46 Ma) with a hiatus from 51.9 to 49.3 Ma (Tauxe et al., 2012). Unit X (Cores 318-U1356A-96R through 100R; 895.5–948.8 mbsf) consists of interbedded stratified and massive sand-
stones, diamicrites, silty claystones, and siltstones. Unit XI (Cores 101R to 106R; 948.8–1000.08 mbsf) consists predominantly of dark green bioturbated claystone and subordinate laminated siltstone and sandstone interbeds (see the “Site U1356” chapter [Expedition 318 Scientists, 2011b]). During the Eocene, Site U1356 was at a latitude of ~65°S, and the dinocyst assemblages suggest a midshelf depositional environment much shallower than today (Pross et al., 2012).

Shore-based analyses of biotic climate proxies (pollen and spores) and indexes based on branched tetra-ether lipids reveals that the climate in lowland settings along the Wilkes Land coast in the early Eocene (Unit XI) supported the growth of highly diverse, near-tropical forests characterized by megathermal floral elements with extremely mild terrestrial winters (>10°C) (Pross et al., 2012), while sea-surface temperatures were extremely warm (Bijl et al., 2013). The middle Eocene (Unit X) is characterized by a marked sea-surface temperature and continental cooling (2°–4°C) (Bijl et al., 2013), loss of megathermal flora, and dominance of a temperate rain forest assemblage (Pross et al., 2012). Furthermore, ongoing work on biomarkers from Site U1356 confirms the presence of the bacterial-derived, C31 (17α, 21β) homohopane within a suite of immature hopanes. The timing of the concentration fluctuations and the molecular structure of hopanes and their compound-specific carbon isotope values (δ13C) reveal the extent to which wetlands fluctuated in response to external, orbital forcing (Toney et al., 2012).

The initial report of alkenones in the Eocene sediments of Site U1356 was significant because alkenones can be utilized both for paleotemperature estimation (e.g., Conte et al., 2006; Prahl and Wakeham, 1987) and as a proxy for the estimation of past pCO2 levels (e.g., Zhang et al., 2013). Thus, we undertook shore-based analysis of 50 samples from Site U1356 to confirm the initial report of the presence of alkenones and to assess whether sufficient concentrations were present for compound-specific δ13C measurements.

**Methodology**

**Shipboard**

Amenable lipids were extracted using a method adapted from Bendle et al. (2007). Full details are given in the “Methods” chapter [Expedition 318 Scientists, 2011a; also see Figure F19 in that chapter]. In brief, sediments were freeze-dried and homogenized, and 2–20 g was hydrolyzed and extracted. The total lipid extract was separated into discrete chemical fractions using methods adapted from Bendle et al. (2007): aliphatic hydrocarbons/n-alkanes (N1), aromatic hydrocarbons (N2), aldehydes and ketones (N3), and alcohols (N4). Alcohols in fraction N4 were derivatized to trimethylsilyl ethers using bis(trimethylsilyl) trifluoroacetamide (BSTFA) reagent. Acidic components were extracted with methylene chloride from the remaining solution after acidifying (pH = 1) with HCl. The acidic fraction was concentrated and the carboxylic acids were derivatized to methyl esters with 14% BF3/methanol (MeOH). The individual lipid fractions were analyzed, and structural identification was confirmed using an HP 5973 gas chromatography mass spectrometer (GC-MS) equipped with a 7683 autosampler and fused silica capillary column (DB-1 60 m × 0.317 mm; internal diameter × 1.50 µm) and mass selective detector (MSD). The GC oven temperature was programmed to ramp at 30°C/min from 50°C to 120°C and then ramp at 5°C/min to 300°C, with a final hold time of 22 min.

**Shore-based extraction and clean-up**

Organic compounds were extracted from 100 powdered and freeze-dried sediment samples at the Royal Netherlands Institute for Sea Research (NIOZ; Utrecht University, Netherlands), and the University of Glasgow (Scotland, United Kingdom) using methods described by Pross et al. (2012). Lipids were extracted by dichloromethane (DCM)/MeOH (9:1, v/v) using the an accelerated solvent extractor (Dionex). The total lipid extract was separated into apolar, ketone, ethyl acetate (EtOAc), and polar fractions over an activated Al2O3 column using hexane: DCM (9:1, v/v), hexane:DCM (1:1, v/v), EtOAc:DCM (1:1 v/v), and DCM:MeOH (1:1, v/v), respectively.

**Ketone fraction GC-MS analyses**

Fifty ketone fraction samples (every other downcore sample from the 100 shore-based extractions) were analyzed using an Agilent GC 7890B coupled to a 5977A MSD GC-MS and equipped with a 7693 autosampler and fused silica capillary column (BP-1 60 m × 0.25 mm internal diameter × 0.25 µm) and mass selective detector at the Birmingham Molecular Climatology Laboratory. The GC oven temperature was programmed to hold for 1 min at 60°C, ramp at 30°C/min from 60°C to 130°C and then ramp at 3°C/min to 345°C, with a final hold time of 15 min. The interface temperature was 340°C. The detector was operated in electron impact mode, scanning from 50 to 800 m/z.
**Results and discussion**

At Site U1356, 25 samples ranging in age from early Eocene to late Miocene were extracted for lipids, separated into chemical fractions, and analyzed by GC-MS onboard the R/V JOIDES Resolution. The compound classes reported include aliphatic hydrocarbons, hopanes, unresolved complex mixtures of branched alkanes, ketones, $n$-alkanols, sterols, diols, and $n$-alkanoic acid methyl esters (FAMES). Relative to the Oligocene and Miocene, molecular concentrations generally increase by several orders of magnitude in the early Eocene sediments. The N1, N3, N4, and FAMES fractions were analyzed by GC-MS, and the results from each fraction are summarized in Tables T10, T11, T12, and T13 in the “Site U1356” chapter [Expedition 318 Scientists, 2011b]), respectively (the N2 and N5 fractions were archived). Examples of GC-MS total- and single-ion chromatograms (of the four analyzed fractions) from four samples are presented in Figures F23 (middle Miocene), F24 (late Oligocene), F25, and F26 (both early Eocene in the “Site U1356” chapter [Expedition 318 Scientists, 2011b]).

Shipboard analysis revealed a grouping of three to four peaks eluting late in the ketone fraction (N3) in >50% of the samples. No alkenone standards were available on ship for comparison. However, based on the general retention time and distribution, these peaks superficially suggested a distribution similar to that expected for the C$_{37:2}$, C$_{38:2}$, and C$_{39:2}$ alkenones. The detector response of these late-eluting peaks was weak when analyzed by the shipboard GC-MS in “scanning mode,” and the resultant mass spectra were inconclusive. However, the late-eluting peaks appeared more prominent when analyzed in “single ion monitoring” (SIM) mode for a number of individual, characteristic mass fragments associated with alkenones (e.g., m/z 81, 109). Thus, the initial shipboard assignation of alkenones was made (see Figs. F25 and F26 and Table T11 in the “Site U1356” chapter [Expedition 318 Scientists, 2011b]).

The additional 50 ketone fraction samples taken from throughout the lowermost 110 m (Units X and XI) of Site U1356 were analyzed on shore to assay for alkenone compounds and confirm the shipboard analyses. As with the shipboard analyses, a grouping of three to four peaks eluted late in the ketone fraction in the majority of the samples. However, the mass spectra of the late-eluting analytes, as produced by shore-based GC-MS, do not show any diagnostic similarity with known alkenone compounds. Shore-based GC-MS results from a representative sample from Site U1356 are given in Figure F1A. For comparison, an alkenone-yielding, Quaternary age sample from the Japan Sea was analyzed in the same laboratory, using the same GC-MS, under the same temperature program and conditions. Figure F1B confirms that positive alkenone identification using the shore-based GC-MS and conditions was feasible, with mass spectra exhibiting characteristic alkenone fragmentation patterns. Moreover, comparison of Figures F1A and F1B unambiguously indicates that the late-eluting peaks in the Site U1356 samples are not alkenones on the basis of both peak retention time and mass spectra. This negative result supersedes the initial report of alkenones in the Eocene sediments of Site U1356 and indicates that shipboard assignation of alkenones was erroneous.

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**References**


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Figure F1. A. Total ion chromatogram of the ketone fraction of an early Eocene sample (318-U1356A-103R-4, 78–77 cm). B. Total ion chromatogram of the ketone fraction of a sample collected from Quaternary sediments from IODP Hole U1425B, Japan Sea. The suites of C_{37}, C_{38}, and C_{39} alkenones are indicated (number in subscript denotes chain length). The molecular structures, specific identifications, and full IUPAC nomenclature are given for the C_{37:2} and C_{37:3} alkenones. Additionally, the electron impact mass spectrogram for the C_{37:2} alkenone is illustrated. See text for details on compound identifications.