PRELIMINARY LIPID ANALYSES OF CORES 49, 54, AND 59 FROM HOLE 462

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ABSTRACT

The solvent-extractable lipids of Sections 462-49-4 (lower Maestrichtian), 462-54-2 (Campanian), and 462-59-1 (Cenomanian), from the Nauru Basin, were investigated for aliphatic and aromatic hydrocarbons, ketones, alcohols, and acids, after separation into acid and neutral fractions. Detailed analyses of the total neutral fraction revealed only n-alkanes, steranes, and triterpanes present in sufficient concentration (>0.5 ng/100 g dry sediment for individual compounds) for component recognition. N-alkanes (C_{19}-C_{35}) constitute the major component of each section, with the most abundant present as 10(n-C_{29}), 135(n-C_{29}), 17(n-C_{26}) ng/100 g dry sediment in Sections 462-49-4, 462-54-2, and 462-59-1, respectively.

The results suggest a high degree of thermal maturity of the lipids, which is interpreted in terms of the high geothermal gradient (about 50°C/km) and the regional volcanicity. The lipid data also suggest that the major source of organic matter was terrestrial and that deposition took place in a low-productivity oxic environment.

INTRODUCTION

The compositions of lipid components from material collected at Site 462 of the Nauru Basin, drilled during Leg 61, have been determined. Three sections have been investigated, by analysis of their extractable lipid components, as two fractions: (1) neutrals (treated with BSTFA to silylate alcohols) and (2) acids as methyl esters (FAME). All sections were low in content of extractable lipids; only n-alkanes, steranes, and triterpanes were detected in sufficient quantities (>0.5 ng/100 g dry sediment) for component recognition.

Table 1 summarizes the lithologic characteristics of cores adjacent to those investigated, and includes dry extracted weights of the samples.

The results are interpreted in an attempt to describe the relationships between the maturity of the sediments and the geological framework. The possibility of pipe dope contamination is also discussed.

EXPERIMENTAL METHODS

The basic experimental scheme for sample analyses and control analyses performed in parallel is shown in Figure 1. It differs from our previous analyses (Barnes et al., 1979; Brassell, Comet, 1980b), in that the small amounts of extractable lipids present, as shown by gas chromatography (GC), precluded subsequent fractionation of the "neutral" fraction by thin-layer chromatography.

GC was performed on a Carlo Erba 2150 gas chromatograph equipped with a 20m OV-1 glass capillary column (0.25 mm i.d.) and He as a carrier gas. The instrument was temperature-programmed at 4°C min⁻¹ from 50 to 275°C, followed by up to 30 minutes isothermal running.

Mass spectra were recorded using a Finnigan 4000 gas chromatograph-mass spectrometer (GC-MS) system equipped with a 20m OV-1 glass capillary column. Data acquisition and processing were controlled by an INCONS 2300 data system. Typical operating conditions were as follows: ion-source temperature, 250°C; electron energy, 35 eV; filament current, 350 µA; scan (m/z 50-500) time, 1 second. The organic carbon values shown in Table 1 were obtained on a Perkin-Elmer 240 CHN analyzer.

RESULTS

General

Inspection of the sediment cores before extraction revealed no visible pipe dope or other shipboard oil-based contamination. Control analyses performed in parallel with sample analyses did not produce detectable amounts of lipid material at similar GC injection volumes.

No alcohols, ketones, or unsaturated hydrocarbons were detected in the neutral lipids by exhaustive mass fragmentographic (MF) searches using key ions (see Brassell, Comet, et al., 1980b; and Brassell, Gowar, et al., in press); only alkanes were detected in all sections. All samples exhibited a "hump" in the GC trace (e.g., Brassell and Eglinton, in press) of the neutral fraction (see Fig. 2). No aromatic hydrocarbons were detected (hence, <0.5 ng/100 g dry sediment), although this may reflect the expected low concentrations of these compounds. No lipids were detected in the "acid" (FAME)
fraction of any section by GC analyses (detection limit about 0.2 ng/100 g).

**Acyclic Components**

$N$-alkanes are the major compound class of each section, ranging from C$_{20}$-C$_{31}$, C$_{19}$-C$_{35}$, and C$_{19}$-C$_{35}$ in Sections 462-49-4, 462-54-2, and 462-59-1, respectively. Their distributions and concentrations (ng/100 g dry sediment) in each section, calculated from GC response, are shown in Figure 3. All samples exhibit a small odd-over-even predominance for their longer-chain $n$-alkanes. Homologues of lower carbon number (<C$_{19}$) were not detected in any section.

No acyclic isoprenoid or simple branched-chain alkanes were detected by MF searches of m/z 85.

**Cyclic Components**

Steranes and diasteranes were detected as minor components in all sections. The small quantities (major component <0.5 ng/100 g dry sediment) in Sections 462-49-4 and 462-59-1, however, precluded characterization of individual compounds. No 4-methyl steroidal hydrocarbons were detected (hence, <0.2 ng/100 g dry sediment).

The major steranes and diasteranes of Section 462-54-2, characterized by MF (m/z 217, m/z 259), mass spectral interpretation, and GC retention time, are listed in Table 2. A partial MF of m/z 217 is shown in Figure 4. Characterization of individual compounds by GC-MS co-injection was precluded by the limited amount of lipid extract. The m/z 217 response of assigned components in Figure 4 may be slightly enhanced owing to coelution of minor components; 13β, 17α-diasteranes, as both 20R and 20S diastereoisomers, are the dominant steroidal hydrocarbons present. Minor 13α, 17β-diasteranes are also present. Identification of individual steranes was complicated by GC coelution with other components and by the low intensity of the mass spectra obtained. Those steranes recognized, however, are tentatively assigned as series of C$_{27}$ to C$_{29}$ 20R and 20S 5α, 14α, 17α-steranes (m/z 149 > m/z 151) and 20R and 20S 5α, 14β, 17β-steranes (m/z 218 > m/z 217) (e.g., Seifert and Moldowan, 1978).

Figure 5 shows the distribution (quantitated from m/z 191) of hopanes that were recognized by their mass spectra, MF of key ions (m/z 149, m/z 177, m/z 191, m/z 205, m/z 219, m/z 233, m/z 247, and m/z 263), and GC retention times. The C$_{29}$-C$_{35}$ hopanes were assigned as 17α, 21β-stereoisomers on the basis of their...
The need to consider pipe-dope contamination as a possibility in such samples (Brassell and Eglinton, in press). Although no evidence of pipe dope was visible in the Site 462 samples, little contamination from such lubricant would be required to dominate the lipid distribution of this similarly organic-carbon-poor suite of samples.

Even though the lipid composition of Section 462-54-2 shows several similarities to that of pipe dope, there are also significant differences (Table 3). From these data—in particular the absence of acyclic isoprenoid alkanes, diterpanes, and aromatic hydrocarbons in Section 462-54-2—we conclude that pipe dope is not a major contributor of lipids to the samples.

Acyclic Components

The n-alkanes form the major compound class of each section. C_{19}–C_{35} n-alkanes and the odd-over-even preference exhibited in the members having higher carbon numbers (most prominent in Section 462-54-2) are characteristic of higher plant waxes (Eglinton and Hamilton, 1967), and therefore suggest a significant terrestrial input to the sediments.

The absence of short-chain n-alkanes, particularly n-C_{17}, a major component of phytoplankton (Oro et al., 1967; Blumer et al., 1971), pristane, and phytane, may reflect low productivity or extensive degradation in an oxic water column. Transport of the terrigenous n-alkane input by clay-mineral adsorption may have screened them from such degradation. Alternatively, the absence of short-chain n-alkanes could suggest loss by the migration of these more volatile components. The absence of squalane and other acyclic isoprenoids suggests that methanogenic bacteria have not been major contributors (cf. Brassell, Comet, et al., in press).

Cyclic Components

The absence of sterols, stanols, and sterenes—which are significant components of immature marine sediments (e.g., Brassell, Comet, et al., 1980b)—together with the presence of steranes, as both the natural 20R and their diagnostically derived 20S isomers (in
approximately a 1:1 ratio) and diasteranes, implies an advanced degree of maturity for the Nauru Basin lipids. Indeed, the stereochemical characteristics of the steranes and diasteranes in Section 462-54-2 resemble those of deeply buried samples (>2400 m), well within the zone of hydrocarbon generation in the Paris Basin (Mackenzie et al., in press). For example, 20R and 20S 24-ethyl-5α,14β,21α-hopanes (IV, R = C15H31; components Q and R in Fig. 4 and Table 2) comprise about 57% of the total 24-ethylcholestanes (components P, Q, R, and S), which is similar to the values for the Courgivaux and Montmirail shales (76% and 73%, respectively) from the Paris Basin (Mackenzie et al., in press). The similar maturity of the steroidal components from the Nauru Basin at a shallower depth of burial (512 m sub-bottom) may, therefore, reflect the regional volcanism that took place between ~110 and ~70 m.y. ago and the resultant high geothermal gradients that affected these Cenomanian to Maestrichtian sediments (Site Summary and Schlanger and Premoli Silva, this volume).

24R-ethylcholeste-5-en-3β-ol is known to be the major sterol of many higher plants, and has been used to infer a terrigenous input to marine sediments (Huang and Meinschein, 1976). The 24-ethylcholestanes and 24-ethylcholestanes, which account for about 60% of the total steroidal hydrocarbons detected in Section 462-54-2, may be derived from 24R-ethylcholeste-5-en-3β-ol (VII), and may therefore reflect a predominantly allochthonous terrestrial input. The absence of 4-methylsteroidal hydrocarbons, which may derive from the 4-methylsterols of various organisms (e.g., dinoflagellates, Brassell, 1980; methanotrophic bacteria, Bird et al., 1971), contrasts with other Cretaceous DSDP sediments (Table 3) and also with the Toarcian shales of the Paris Basin (Ensminger, 1977).

The series of 17α,21β-hopanes (VI), as both the natural 22R and their diagenetic 22S epimers, appear at a late stage in the maturation of sedimentary hopanes (Van Dorsselaer et al., 1977; Ensminger, 1977). 17α, 21β-hopanes are believed to be derived from their 17β, 21β counterparts by a clay-catalyzed diagenetic isomerization of the hopane compounds synthesized by bacteria (De Rosa et al., 1973; Rohmer and Ourisson, 1976); 17β, 21β-hopanes are entirely isomerized to their more stable counterparts above the zone of hydrocarbon generation (Ensminger et al., 1977; Mackenzie et al., in press). Their absence in these samples is consistent with a mature sedimentary hopane distribution. The 22S diastereoisomers comprise about 57% of the total extended (>C31) 17α,21β-hopanes (Appendix); this value is similar to the values (50-65%) for deeper samples (>2000 m) from the Paris Basin (Ensminger et al., 1977; Mackenzie et al., in press). In the Nauru Basin, the maturity indicated by the hopane distributions is consistent with that inferred from the stereochemistry of the steroidal hydrocarbons.

Various organisms, including algae, ferns, and bacteria (Ensminger, 1977), are possible sources of C30 hopanes and degraded (<C30) hopanes. However, extended hopanes are thought to derive uniquely from the polyhydroxybacteriohopanes (Rohmer and Ourisson, 1976; Ourisson et al., 1979), and therefore suggest a bacterial input to the sediments.

Influence of the Underlying Volcanic Complex

Previous reports of thermal effects of intrusive basalt sills in the Kimmeridgian shales, Milnes Island, Greenland (Perregrand and Schiener, 1979) and Cape Verde Islands (Dow, 1978) suggest dramatic thermal alteration of organic matter within the proximity of roughly half the thickness of a sill, and a rapid decrease in such effects at increasing distance from the sill. Section 462-59-1 lies 8 meters above the shallowest intrusive basalt sill, but comparison of the distributions of n-alkanes and hopanes common to all sections indicates that Section 462-59-1 has not experienced a significantly more severe thermal history than the two overlying samples. It appears, therefore, that the thermal effects of the volcanic complex are not markedly different throughout the sedimentary sequence examined herein (462-551 m sub-bottom depth; 97-8 m above the shallowest sill). The full extent of sediment maturation effected by the volcanic complex, in particular its influence on shallower horizons, cannot be fully assessed from the avail-
TABLE 3. Semi-quantitative comparison of Section 462-54-2 with two other Cretaceous samples
and with pipe dope.

<table>
<thead>
<tr>
<th>Location</th>
<th>Chronostratigraphy</th>
<th>Lithology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nauru Basin</td>
<td>Campanian</td>
<td>Marly limestone</td>
</tr>
<tr>
<td>Bay of Biscay</td>
<td>Lower Albian</td>
<td>Carbonaceous marly limestone</td>
</tr>
<tr>
<td>Moroccan Basin</td>
<td>Valanginian</td>
<td>Turbidite</td>
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</table>

<table>
<thead>
<tr>
<th>Section:</th>
<th>462-54-2</th>
<th>402A-24-1</th>
<th>416A-18-2</th>
<th>Pipe dope</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound Class</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N-alkanes</td>
<td>+ + +</td>
<td>+ + +</td>
<td>+ + +</td>
<td>+ +</td>
</tr>
<tr>
<td>CPI of n-alkanes</td>
<td>(CPI 1.3)</td>
<td>(CPI 1.4)</td>
<td>(CPI 2.4)</td>
<td></td>
</tr>
<tr>
<td>Acyclic isoprenoids</td>
<td>n.d.</td>
<td>+ + +</td>
<td>+ + +</td>
<td>+ +</td>
</tr>
<tr>
<td>Extended cyclic diterpanes</td>
<td>n.d.</td>
<td>+ +</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>Diasterenes</td>
<td>n.d.</td>
<td>+ +</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>Steranes</td>
<td>+ +</td>
<td>+ +</td>
<td>+ +</td>
<td>+ +</td>
</tr>
<tr>
<td>Steranes</td>
<td>+ +</td>
<td>n.d.</td>
<td>+ +</td>
<td>n.d.</td>
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<tr>
<td>Diasterenes</td>
<td>+ +</td>
<td>n.d.</td>
<td>+ +</td>
<td>n.d.</td>
</tr>
<tr>
<td>4-Methylsteroidal hydrocarbons</td>
<td>n.d.</td>
<td>+ +</td>
<td>+ +</td>
<td>+ +</td>
</tr>
<tr>
<td>4α-Hopanes</td>
<td>+ +</td>
<td>n.d.</td>
<td>+ +</td>
<td>+ +</td>
</tr>
<tr>
<td>Aromatic hydrocarbons</td>
<td>+ +</td>
<td>+ +</td>
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</tr>
<tr>
<td>“Hump”</td>
<td>+ +</td>
<td>+ +</td>
<td>+ +</td>
<td></td>
</tr>
</tbody>
</table>

+ + + major 10-100%; + + minor 1-10%; + trace <1%, of major component, respectively.
1 n.d. = not detected.
2 Data presented herein.
4 Brassell, Comet, et al., in press (b).
5 Brassell and Eglinton, in press.
6 Range C22–C35.
7 Range C36–C34.

Table 3 illustrates the overall thickness of the mid-Cretaceous volcanic complex (>500 m; Site Summary, this volume) suggests, however, that such alteration is probably extensive.

**Input Considerations**

An inconsistency in the neutral lipid fraction is the coexistence of an alkane "hump" and an odd-over-even preference in the long-chain n-alkanes. Such a combination of lipid characteristics suggests that they are not wholly indigenous but contain, in part, migrated components. The relative contributions of different sources to the lipid distribution are uncertain, but a significant terrestrial source would seem to be indicated, with little evidence of an "algal" component.

**Comparison of Nauru Basin Lipids with Those of Other Cretaceous DSDP Samples**

The small amounts of lipids in Section 462-54-2 preclude detailed quantitative comparisons with Leg 48 and 50 samples; some comments can, however, be made (Table 3).

The lipids of the Nauru Basin are markedly more mature than those of the other cited samples, as indicated by the following:

1) CPI of n-alkanes.
2) Absence of alkenes (e.g., diasterenes, hopanes).
3) Sterane stereochemistry.
4) Predominance of 17α,21β-hopanes over 17β, 21β-hopanes.
5) Absence of functionalyzed lipids (e.g., alkanols, alkanones, and alkanolic acids) found as minor components of the Leg 48 and Leg 50 samples (Wardroper, 1979; Brassell, 1980).

In summary, the Nauru Basin lipids are characteristic of mature sediments, in contrast to lipids of samples from the eastern North Atlantic, where immature lipid distributions have been recognized.

**CONCLUSIONS**

The three Upper Cretaceous samples contain low concentrations of organic material of mainly terrestrial origin. The apparent absence of organic material attributable to algal sources and the low organic carbon values probably reflect low productivity and oxic conditions in the water column. The predominance of hopanes over other polycyclic alkanes may imply extensive microbial activity and reworking of the organic matter. It is possible that the lipids are not wholly indigenous and may, in part, include migrated components.

Deposition of these Nauru Basin sediments occurred during the emplacement and growth of an off-ridge volcanic sill complex, but regional heating was never sufficient to obliterate the “magnetic stripe pattern” (Site Summary, this volume). The organic matter does, however, reflect this thermal episode. The lipid extracts are atypically mature for such shallow depths of burial (462–551 m sub-bottom), apparently lying within the oil window. Oil generation could therefore occur within the Nauru Basin, although it would be expected to be limited, owing to the small amounts of organic matter present in these samples. These data may be explained in terms of high paleotemperatures, although interpretation is complicated by the possible presence of migrated lipids.

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