11. BIOMARKER AND PYROLYSIS GEOCHEMISTRY OF ORGANIC MATTER FROM ARGO AND GASCOYNE ABYSSAL PLAIN SEDIMENTS, NORTHEASTERN INDIAN OCEAN

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ABSTRACT

The sediments of the Argo and Gascoyne abyssal plains are generally lean in organic matter, are immature, and contain hydrocarbons trapped during sediment deposition rather than those generated during sediment catagenesis. TOC concentrations in the Argo Abyssal Plain Cenozoic sediments are 0.5 wt%, and organic matter appears to be from mixed marine and reworked, degraded, organic matter sources, with the latter being contributed by turbidity flows from the nearby continental margin. TOC concentrations within the Cenozoic sediments of the Gascoyne Abyssal Plain are mostly undetectable (<0.1 wt%). Biomarker distributions determined by gas chromatography (GC) and gas chromatography-mass spectrometry (GC-MS) indicate that organic matter extracted from the Lower Cretaceous sediments from both sites is predominantly marine with varying contributions from terrestrial organic matter. The specific marine biomarker, 24-n-propylolestanate is in relatively high abundance in all samples. In addition, the relatively high abundance of the 4-methylsteranes with the 23, 24-dimethyl side chain (in all samples) indicates significant dinoflagellate contributions and marine organic matter.

The ratios of \( n-C_{28}/n-C_{17} \) reflect relative contributions of marine vs. terrestrial organic matter. TOC, while generally low at Argo, is relatively high near the Barremian/Aptian boundary (one sample has a TOC of 5.1 wt%) and the Aptian/Albian boundary (up to 1.3 wt% TOC), and two samples from the Barremian and Aptian sections contain relatively high proportions of terrestrial organic carbon. TOC values in the Lower Cretaceous sediments from Gascoyne Abyssal Plain are low (<0.1 wt%) near the Aptian/Barremian boundary. TOC values are higher in older sediments, with maxima in the upper Barremian (1.02 wt%), the Barremian/Hauterivian (0.6 wt%), and Valanginian (1.8 wt%). Sediments from the upper Barremian contain higher amounts of terrestrial organic carbon than older sediments.

INTRODUCTION

The formation of the modern Indian Ocean, including the Argo and Gascoyne Abyssal Plains, off northwestern Australia (Fig. 1), began with Late Jurassic to Early Cretaceous rifting along the northern and western margins (Ludden, Gradstein et al., 1990). Geohistory analysis of the nearby DSDP Site 261 in the Argo Abyssal Plain indicates an initial water depth of around 2.5 km. Rapid cooling and subsidence resulted in a water depth near 4000 m during the Hauterivian/Barremian. Slower subsidence through the Late Cretaceous and Cenozoic resulted in a present-day water depth near 5700 m (Ludden, Gradstein et al., 1990). The Gascoyne Abyssal Plain (Site 766, Fig. 1) is near the foot of the Exmouth Plateau escarpment. Subsidence and sedimentation considerations indicate this site originated at a water depth near 800 m; rapid subsidence in the Early Cretaceous brought the site near to its present water depth of 4000 m.

The sediments at the Argo and Gascoyne abyssal plains began to be deposited during the Early Cretaceous with the onset of seafloor spreading. About 935 m of sediment was cored at Site 765 (Argo), and approximately 500 m of these sediments are Lower Cretaceous calcareous claystones and hemipelagic clays that accumulated at rates of about 5 meters per million years (m/Ma) during Valanginian through Barremian time, 14 m/Ma during the Aptian, and 4.2 m/Ma during the Albian (Ludden, Gradstein, et al., 1990). Site 765 has remained close to or below the CCD. Some turbidite deposition is evident in the Lower Cretaceous sequence. Late Cretaceous and Paleogene sedimentation rates were low (<2 m/Ma), and the appearance of siliciclasti-
Figure 1. Map of the continental margin of northwestern Australia showing drill Sites 765 (Argo Abyssal Plain) and 766 (Gascoyne Abyssal Plain).

<table>
<thead>
<tr>
<th>Core, section, interval, (cm)</th>
<th>Age</th>
<th>TOC (wt%)</th>
<th>$T_{\text{max}}$ (°C)</th>
<th>$S_1$</th>
<th>$S_2$</th>
<th>PI</th>
<th>HI</th>
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<td>765B-03L-3, 8–16 P</td>
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<td>545</td>
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<td>0.00</td>
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<td>0.00</td>
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</tr>
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<td>765C-44R-3, 3–50 Ap</td>
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<td>417</td>
<td>0.19</td>
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<td>0.13</td>
<td>100</td>
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<td>765C-44R-5, 2–10 Ap</td>
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<td>0.00</td>
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<tr>
<td>765C-59R-1, 96–104 B</td>
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<td>354</td>
<td>0.07</td>
<td>0.27</td>
<td>0.21</td>
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<td>0.00</td>
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<tr>
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<td>414</td>
<td>0.04</td>
<td>0.61</td>
<td>0.06</td>
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<tr>
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<tr>
<td>765C-32R-4, 132–138 B/H</td>
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<td>75</td>
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<td>334</td>
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<td>0.17</td>
<td>130</td>
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<tr>
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<td>362</td>
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<tr>
<td>765C-37R-1, 59–65 H</td>
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<td>72</td>
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<td>0.34</td>
<td>109</td>
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</tr>
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<td>0.33</td>
<td>0.58</td>
<td>0.37</td>
<td>132</td>
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</tr>
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<td>0.57</td>
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<tr>
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<td>371</td>
<td>0.22</td>
<td>0.54</td>
<td>0.29</td>
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<tr>
<td>765C-43R-4, 114–119 H</td>
<td>0.53</td>
<td>377</td>
<td>0.21</td>
<td>0.59</td>
<td>0.26</td>
<td>111</td>
<td></td>
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<tr>
<td>765C-44R-4, 98–106 H</td>
<td>1.25</td>
<td>424</td>
<td>0.33</td>
<td>1.27</td>
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<tr>
<td>765C-46R-6, 50–57 V</td>
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<td>0.61</td>
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<td>84</td>
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</tr>
<tr>
<td>765C-47R-4, 95–102 V</td>
<td>0.68</td>
<td>403</td>
<td>0.21</td>
<td>0.65</td>
<td>0.24</td>
<td>96</td>
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</tr>
<tr>
<td>765C-48R-4, 16–18 V</td>
<td>0.50</td>
<td>395</td>
<td>0.15</td>
<td>0.70</td>
<td>0.18</td>
<td>140</td>
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</tr>
<tr>
<td>765C-49R-3, 47–51 V</td>
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<td>399</td>
<td>0.27</td>
<td>0.88</td>
<td>0.24</td>
<td>124</td>
<td></td>
</tr>
</tbody>
</table>

Age: P = Pleistocene, Ap = Aptian, B = Barremian, B/H = Barremian/Hauterivian, V = Valanginian. $S_1$, $S_2$ in mg HC/g; PI and HI in mg HC/g organic C.
Bon fractions were then analyzed for a variety of biomarkers by combined gas chromatography-mass spectrometry (Summons et al., 1988).

TOTAL ORGANIC CARBON DISTRIBUTIONS, PYROLYSIS, AND SOURCE CHARACTERISTICS

Argo Abyssal Plain

Shipboard total organic carbon (TOC) data from the Argo Abyssal Plain are summarized in Figure 2. Four additional samples were selected from the Lower Cretaceous sediments of Site 765 of Aptian through Barremian age, while one sample was taken from the Pleistocene section (BMR Rock-Eval data shown in open circles). The TOC values from these additional analyses (Table 1) vary between 0.4 and 1.3 wt% and are similar to those found from the shipboard analyses.

Argo Abyssal Plain (Cenozoic Sediments)

TOC data from the Cenozoic sediments are highest in the top 50 mbsf (up to 1.5 wt%), and decrease with increasing depth, although the data show large concentration differences over short distances. These sediments are dominated by turbidite deposition (Ludden, Gradstein, et al., 1990). Calcium carbonate and TOC data from a single turbidite are shown in Figure 3. Calcium carbonate data are lowest in the top of the turbidite, probably because of dissolution in corrosive bottom waters below the CCD. TOC concentrations vary about tenfold over about 1.5 m of this turbidite and are lowest at the base of the turbidite in the coarse-grained sediments and highest in the fine-grained sediments toward the turbidite top. These large variations within a single turbidite, over distances of less than 2 m, are comparable to changes in TOC over the whole Cenozoic section and thus explain, in part, TOC variations in the Cenozoic sediments.

Pore water data from this site (Ludden, Gradstein, et al., 1990) show that sulfate decreases with increasing depth in the sediments to near asymptotic concentrations of 6 to 7 mM, representing about a 70% depletion from bottom-water concentrations, at about 300 mbsf. The oxidation and remineralization of buried organic matter, primarily by seawater metabolites, oxygen, nitrate, and sulfate during early diagenesis (e.g., Bender and Heggie, 1984) explains the trend of decreasing TOC with increasing depth of burial. The observed TOC distributions with depth in the Cenozoic sediments of the Argo Abyssal Plain thus result from the combined effects of pelagic and frequent turbidite deposition with a diagenetic overprint.

Rock-Eval Data

The combined Rock-Eval data sets from Site 765 were screened, and those having S2 <0.2 and TOC <0.5 wt% were not considered further (the screened Rock-Eval data from shipboard analyses are summarized in Table 2). A cross plot (Peters, 1986) of the hydrogen index (HI) and oxygen index (OI) of Site 765 data is shown in Figure 4. This plot may be used similarly to a van Krevelen diagram to differentiate organic matter types. Because all samples have TOC <2 wt% (with one exception) and have variable amounts of calcium carbonate and clay minerals, the pyrolysis yields have probably been suppressed by the mineral matrix effect (Espitalié et al., 1980; Orr, 1983; Katz, 1983; Peters, 1986). This effect results in low, but variable, HI values compared to results on demineralized organic matter and, hence, introduces artificial differences in organic matter compositions.

The Cenozoic (Fig. 4) samples were from the top 100 mbsf of sediment and are of Pleistocene age. These samples have variable
HI values (150<HI<500) and plot between the Type II/III mean evolution lines (data not plotted have HI<400 and OI>350), suggesting predominantly Type II organic matter, with the variation in HI values probably owing to a variable mineral matrix effect. An additional contribution from varying amounts of hydrogen-poor, reworked organic matter is also likely (see discussion in “Maturity” section, this chapter). Shipboard palynological investigations indicated that samples from Miocene/Pliocene mass-flow deposits contained reworked Jurassic dinoflagellates, as well as spores and pollen (Ludden, Gradstein, et al., 1990). The variation in OI is more problematic. For example, in the Cretaceous marine Toolebuc Formation (Queensland, Australia), which is Type II organic matter (Boreham and Powell, 1987), oxidation (of the marine organic matter) results in a decrease in HI with little increase in OI, whereas high OI and low HI values were associated with terrestrial organic matter. The trend in the Cenozoic samples in Figure 8 (decreasing HI with increasing OI) also suggests a combination of varying degrees of oxidation associated with mixed marine and terrestrial sources of organic matter. However, in these organic-lean rocks, a major contribution to the amount of CO₂ released is most likely from decomposition of carbonate minerals, which would mask any trend in the data.

**Argo Abyssal Plain (Mesozoic Sediments)**

TOC are generally low (wt%) in the Mesozoic section. However, isolated samples of higher TOC were occasionally found. The highest TOC was found in an isolated, thin (1 cm), black sediment layer (123-765C-45R-4, 100-101 cm) near the Barremian/Aptian boundary (Fig. 2). This sediment (Thurow, pers. comm., 1991) was finely laminated and not bioturbated (although adjacent sediments were bioturbated) and was (under the scanning electron microscope) almost entirely framboidal pyrite. These observations and other estimates of the pyrite abundance (Reggie, this volume) suggest that this sediment was deposited during a brief period of anoxia in the overlying water (additional samples for laboratory analyses, however, were not available from this interval). TOC values near the Aptian/Albian boundary were up to 1.3 wt% (Fig. 2).

The Mesozoic samples of Site 765, from depths greater than 650 mbsf and representing Lower Cretaceous sediments from Albian through Valanginian/Berriasian ages, have a low HI (75) and cluster about the Type III mean evolution line. These data

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**Table 2. Selected shipboard Rock-Eval and calcium carbonate data, Site 765.**

<table>
<thead>
<tr>
<th>Core, section, interval (cm)</th>
<th>Depth (mbsf)</th>
<th>CaCO₃ (wt%)</th>
<th>S₁</th>
<th>S₂</th>
<th>TOC (wt%)</th>
<th>C</th>
<th>HI</th>
<th>OI</th>
<th>T_max (°C)</th>
<th>PI</th>
<th>S₂/S₁</th>
</tr>
</thead>
<tbody>
<tr>
<td>765B-1H-1, 96-98</td>
<td>0.96</td>
<td>13.0</td>
<td>0.64</td>
<td>1.47</td>
<td>4.98</td>
<td>0.85</td>
<td>0.17</td>
<td>172</td>
<td>585</td>
<td>384</td>
<td>0.30</td>
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<tr>
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<td>52.5</td>
<td>0.38</td>
<td>1.25</td>
<td>5.15</td>
<td>0.52</td>
<td>0.13</td>
<td>240</td>
<td>990</td>
<td>404</td>
<td>0.23</td>
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<tr>
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<td>0.68</td>
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<td>0.28</td>
<td>452</td>
<td>172</td>
<td>490</td>
<td>0.20</td>
</tr>
<tr>
<td>765B-1H-6, 61-63</td>
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<tr>
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<td>3.61</td>
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<td>401</td>
<td>548</td>
<td>0.16</td>
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</tbody>
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**Figure 3. Calcium carbonate and TOC data from a turbidite.**
suggest that these samples have a high content of terrestrial organic matter, although the mineral matrix effect (because of the relatively low calcium carbonate and relatively high clay contents) probably results in depressed HI compared to the Cenozoic samples. The TOC-rich sample (5.1 wt%) had the highest HI, but the Rock-Eval data from this and adjacent samples did not provide any clues to changes in the type of organic matter being deposited in the sediments.

Gascoyne Abyssal Plain

Shipboard TOC data in sediments from the Gascoyne Abyssal Plain are summarized in Figure 5. TOC data from 25 additional samples (open circles) selected from the Lower Cretaceous (Table I) were similar to shipboard data. TOC was mostly undetectable in the Cenozoic and Upper Cretaceous sediments, but increased in the Lower Cretaceous to concentrations of 1.8 wt% in samples of Valanginian age. The Rock-Eval data determined aboard the Resolution were screened, and all samples having S<sub>2</sub> < 0.2 and TOC < 0.5 wt% have been excluded—the remainder are summarized in Table 3. The Mesozoic samples (open triangles) are plotted in Figure 4. All Cenozoic samples failed to provide reliable Rock-Eval parameters and have been excluded. The Cretaceous samples have HI values of generally < 100 and variable OI, and contain Type III organic matter similar to that of the Lower Cretaceous sediments from Argo.

GAS CHROMATOGRAPHY AND SOURCE CHARACTERISTICS

Gas chromatography (GC) analyses of the saturates' extract of all samples showed a homologous series of straight-chained alkanes ranging from n-C<sub>15</sub> to >n-C<sub>30</sub>. Typical GC traces from Early Cretaceous samples (of Aptian age) from Argo (i.e., Sample 123-765C-44R-3, 42–50 cm) and a Valanginian sample from Gascoyne (123-766A-48R-4, 16–18 cm) are shown, respectively, in Figures 6A and 6B. These traces represent the end-member compositions and illustrate the differences between the characteristics of the saturated hydrocarbons at each of these two sites.

Three important features are found in the gas chromatograph (FID) traces. The first relates to the relative proportions of short- and long-chain n-alkanes summarized in Table 4. The ratio of n-C<sub>27</sub>/n-C<sub>17</sub> may be used as an indicator of the relative contributions of terrestrial and marine organic matter respectively (e.g., Tissot and Welte, 1984). The high n-C<sub>27</sub>/n-C<sub>17</sub> ratios measured in the relatively thick (115 m) Aptian (Sample 123-765C-44R-3, 42–50 cm) and Barremian (85 m) sections (Sample 123-765C-49R-1, 96–104 cm) from Site 765 indicate high proportions of terrestrial organic matter. Similarly, the high ratios measured in two samples from Site 766 (123-766A-30R-4, 48–54 cm and 123-766-31R-2, 145–150 cm) in dark greenish-gray to black upper Hauterivian and Barremian calcareous claystones indicate a higher proportion of terrestrial organic matter in these samples than in older sediments from Site 766. These observations are consistent with palynological shipboard investigations, which found an abundance of spores and pollen in the high sedimentation rate (14 m/m.y.) Aptian/Albian sections of Site 765 and also in the M. Australis Zone (mostly of Barremian age) of Site 766 (Ludden, Gradstein, et al., 1990).

The second feature of the chromatograms (Fig. 6) is the high proportion of peaks eluting between each alternative n-alkane (peaks indicated by an asterisk). Their elution positions and mass spectra suggest that these are homologous series of branched alkanes, but at present, little is known about either their precise structure or their origin. These branched alkanes have different characteristics from the mid-chain monomethyl alkanes that are prominent in many Proterozoic and lower Paleozoic sediments.
and petroleums (Jackson et al., 1986; Klomp, 1986; Summons, 1987; Summons et al., 1988). These unidentified hydrocarbons sometimes appear in samples of sedimentary bitumen that have experienced high degrees of thermal alteration (Summons, unpubl. data, 1990), or in immature, organically lean, oceanic sediments. The former, however, is not consistent with other indicators of the low thermal maturity of samples from both sites. Here, their presence, along with an extensive unresolved complex multiple (UCM) of hydrocarbons, is consistent with a suite of structures that no longer resemble their original biogenic precursors and that might indicate a component of reworked or “background” organic matter. Gough and Rowland (1990) showed by chemical degradation that the UCM (hump) owes its appearance to a multiplicity of unnaturally branched structures. Their high relative

![Figure 6. Representative GC traces of (A) a sample of Aptian age from Site 765 and (B) a sample of Valanginian age from Site 766.](image-url)
abundance here would result from the fact that little or no hydrocarbon has been generated by catagenesis of in-situ kerogen.

Finally, the two major acyclic isoprenoid hydrocarbons in all the samples are pristane and phytane. Empirical observations of marginally mature to mature rocks suggest that the ratios of pristane to phytane vary systematically according to the type of depositional environment (Powell and McKirdy, 1973). However, the exact reasons for this are currently subject to some controversy (see reviews by Volkman and Maxwell, 1986; ten Haven et al., 1985, 1987, 1988). The sources of pristane and phytane are multiple and complex, but as a general rule, Pr/Ph values >3 are typical of organic matter having predominantly terrestrial sources deposited in, or transported through, an oxic environment. Values of less than 1 are typically found in sediments from anoxic marine or hypersaline depositional environments. Because pristane/phytane ratios are only reliable indicators of source environment in marginally mature to mature rocks, the immaturity of all Mesozoic samples from both sites (see below) and the fact that Pr/Ph ratios mostly fall between 1 and 3 preclude a definitive interpretation of the ratios summarized in Table 4 and, therefore, of depositional environment and occurrences of anoxic/oxic conditions (Powell, 1988).

Interpretations of biomarker ratios, such as Pr/Ph and others (see below), to infer the depositional setting and sources of organic carbon must be made cautiously and in consideration of other known information. In this case, the indicators of low maturity tell us that only a minor proportion of the organic carbon has been converted to hydrocarbons. As such, these hydrocarbons may not be representative of the whole suite of organic matter types, because thermally stable kerogen will only release diagnostic markers at higher levels of maturation. Hence, the source-related biomarkers that we identified can indicate that contributions of different types exist, but cannot be used to define accurately the quantitative relationships of these contributions.

**GC-MS and Source-Related Biomarkers**

Information regarding source type sometimes may be obtained from the relative abundance of C_{27}, C_{29} sterols or their fossil counterparts, the steranes (Huang and Meinschein, 1979; Shi et al., 1982). Land plant influxes usually result in a dominance of C_{29} desmethylsteranes, whereas in predominantly marine sediments, different types and proportions of algal to zooplankton sources can result in a wide variety of sterane distributions (Seifert and Moldovan, 1978; Mackenzie et al., 1982; Volkman, 1986). Geological age also may be an important consideration (Granholm, 1986; Granholm and Wakefield, 1988) when assessing the C_{27}/C_{29} parameter. The lower C_{27}/C_{29} ratios for the Barremian/Hauterivian through Valanginian sediments of Site 766, compared with the Aptian and Barremian sediments of Site 765 (Table 5, Col. 1), suggest an increased terrestrial influence in the Valanginian sediments of Site 766. However, this is contrary to the n-alkane distribution (Table 4) and highlights the difficulties for attributing specific sources to the desmethylsteranes (Volkman, 1988). In this case, we suggest that the n-alkane distributions better reflect the terrestrial influxes. The presence and high relative abundance of the specific C_{35} sterane, 24-n-propyloleoholostane, is a reliable marine marker (Moldowan et al., 1985, 1990). Its relative abundances here (Table 5, Col. 2) are similar, for example, to the Mesozoic marine Toolebuc Formation (Boreham and Powell, 1987) and other classic marine sediments, such as the Monterey Formation of California, Lower Toarcian of France, and North Sea Kimmeridge Oil Shale (Moldowan et al., 1985, 1990; Summons et al., 1987) and, consequently, indicate a major contribution of organic matter of marine origin to the Argo and Gascoyne abyssal plains.

Methylsteranes are in high abundance relative to their desmethyl counterparts in all samples (Table 5, Col. 3). The 4-methyl substitution in ring-A is a characteristic of dinoflagellate sterols and suggests that these organisms are the source of the 4-methylsteranes (Whithers, 1983; Rubinstein and Albrecht, 1975; Robinson et al., 1984; Brassell et al., 1987). The presence of the steroid side chain substitution of 24-ethyl, together with four isomers of 23,24 dimethyl for the C_{35} 4-methylsteranes (Table 5; Col. 4), is a characteristic of marine sediments (Summons et al., 1987). This pattern becomes prominent in the middle Triassic and coexists with the first appearance in marine sediments of dinoflagellate cysts (Thomas et al., 1990). In nonmarine sediments of equivalent age, only the 24-ethyl side-chain substitution pattern is found in 4-methylsteranes (Summons et al., 1987). The higher relative abundance of the 24-ethyl to the 23,24 dimethyl side chain (dinosterane) (Table 5; Col. 4) in the Aptian and upper Barremian sediments from Argo may reflect subtle compositional differences between dinoflagellate lipid influxes. Two additional series of methylsteranes, 2α- and 3β-methyl ring-A substitution, were observed together in near equal concentrations as their 4-methyl isomers (Table 5; Col. 5). Some evidence exists to suggest that the occurrence of 2- and 3-methylsteranes is associated with bacterial activity during early diagenesis (Summons and Capon, 1988). This appears to be true for the present sample set. The Aptian and upper Barremian sediments from Argo, which

| Table 5. Gas chromatography and mass spectrometry source-dependent biomarker parameters. |
|------------------------------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|
| Core, section, interval (cm)            | C_{27} Sterane   | C_{29} Sterane   | C_{30} Me Sterane| C_{30} Me Sterane| C_{29} Sterane   | C_{30} Me Sterane| C_{30} Me Sterane| C_{30} Me Sterane|
|                                         |                  |                  |                  |                  |                  |                  |                  |                  |
| 765C-4R-5, 2-10                          | 0.84             | 0.087            | 0.51             | 0.90             | 1.60             | 5.5              | 0.02            | 0.51             | 0.024            | 1.1              |
| 765C-4R-1, 96-104                        | 1.10             | 0.070            | 0.32             | 0.48             | 1.21             | 13.3             | 0.032           | 0.72             | 0.023            | 2.0              |
| 766A-33R-3, 98-104                       | 0.53             | 0.087            | 0.32             | 0.48             | 0.77             | 3.5              | 0.016           | 0.80             | 0.051            | 2.9              |
| 766A-45R-5, 9-106                        | 0.52             | 0.120            | 0.44             | 0.48             | 0.29             | 2.3              | 0.027           | 0.27             | 0.034            | 2.2              |
| 766A-7R-4, 95-102                        | 0.46             | 0.065            | 0.37             | 0.63             | 0.48             | 2.9              | 0.034           | 0.47             | 0.047            | 3.5              |
| 766A-8R-14, 16-18                        | 0.62             | 0.065            | 0.45             | 0.33             | 0.57             | 3.8              | 0.041           | 0.45             | 0.048            | 2.2              |
| 766A-49R-3, 47-51                        | 0.65             | 0.110            | 0.48             | 0.58             | 0.53             | 4.1              | 0.029           | 0.51             | 0.044            | 3.0              |
have the highest relative concentration of hopanes, (Table 5; Col.
6, an indicator of bacterial activity, see below), also contain more
abundant 2- and 3-methylsteranes relative to 4-methylsteranes
(Table 5; Col. 5).

A significant and variable bacterial contribution can be seen from
the relative abundance of hopanes to steranes (Table 5; Col.
6). The Aptian and upper Barremian samples from Argo contain
the highest concentration of hopanes relative to the desmethyl-
steranes. Sedimentary hopanes are known to be derived from
functionalyzed hopanoids in bacteria (Rohmer et al., 1984; Ouris-
on et al., 1987), while hopanoids and hopane polysaccharides having an
additional methyl substitution at C-2 or C-3 in ring-A have been
recognized in several classes of bacteria, such as methylocroplastic
bacteria and cyanobacteria (Ourisson et al., 1987). Price et al.
(1988) and Summons and Jahnke (1990) observed that high rela-
tive abundances of methylhopanes (i.e., 2-methylhopanes) in the
range 10% to 20% of corresponding hopane are characteristic of
sediments having high carbonate contents. The relative abun-
dances of 2-methylhopanes in the samples from both sites (Table
5; Col. 7) are low and in the range 1.2% to 4.1% of the correspond-
ing hopanes. This is consistent with the low carbonate contents,
and the ratio of 2-methylhopane/hopane does not vary signifi-
cantly over the sample set. The ratio of C29 hopanes/C30 hopanes
(Table 5, Col. 8) is also typical of clastic sediments.

Other distinctive hopanes include the C28 species, 28,30-bis-
norhopane, commonly found in marine sediments (e.g., Grantham
et al., 1980; Curiale and Odermatt, 1989) and 29,30-bisnorhopane
(Summons and Powell, 1987). Both these biomarkers are present
in low abundances in all samples (Table 5; Cols. 9 and 10) but
their presence, in accord with other source parameters, suggests
a marine source for the organic matter. The high relative abun-
dances of bacterial biomarkers is a further illustration of the
importance of bacterial contributions to sedimentary organic
carbon irrespective of the type and age of the depositional setting.

MATURITY

Pyrolysis data are summarized in a plot of T_max vs. HI (Fig. 7).
The Cenozoic samples appear to separate into two distinct groups:
those samples with T_max <430°C and those greater than 450°C.
The former T_max values suggest that the samples are immature,
while the latter indicate mature to over-mature organic matter.
However, based upon these limited data, one can see that two
carbon sources are contributing to the TOC in these organic-lean
sediments. A high proportion of reworked allochthonous organic
matter will manifest itself as a high T_max value, while the immature
autochthonous component is responsible for the consistently high
HI values. Where this latter component is high, a low T_max also is
observed. Based on the T_max values, the organic matter in the
Mesozoic samples from both sites is immature.

GC analyses of the saturated hydrocarbons indicate that a
pronounced odd-to-even predominance exists in the hydrocar-
bons greater than C28 in all samples, reflecting the immaturity of
the sediment with respect to petroleum generation (Tissot and
Welte, 1984). The high proportions of moretane 17β,21α (>-10%)
compared to hopane (17α,21β) (Table 6; Col. 3), and the
dominance of Tm over the more stable Ts (Table 6; Col. 4)
constitute further evidence of the immature nature of the sedimen-
ta's maturity. The relative maturity within the sample set is harder to ascer-
tain because these maturation parameters are also influenced to
varying degrees by source effects. However, it appears that no
significant maturity difference exists among samples from the two
sites, although petrography of the organic matter (which was not
done) might clarify the question of maturity.

SUMMARY

Examination of GC, GC-MS biomarker, and Rock-Eval data
derived from samples analyzed at BMR and Rock-Eval data
gathered aboard the Resolution indicates the following about the
nature and source of organic matter at these two abyssal plain drill
sites:

1. Cenozoic sediments from Site 766 have undetectable TOC
(<0.1 wt%). Cenozoic sediments from Argo have <1.5 wt% TOC,
and concentrations tend to decrease downhole (although erratically because of turbidite deposition) as organic carbon is remini-eralized via early diagenetic reactions.

2. Rock-Eval pyrolysis of Cenozoic samples from Argo suggests that the organic matter is mostly of a marine source, but with a component of reworked and degraded organic matter, probably derived from the continental margin via turbidite deposition.

3. Biomarker distributions determined by GC and GC-MS analysis indicate that the organic matter extracted from the sediments (inferred from the high relative abundances of the specific marine biomarker, 24-n-propynoylechsterol), at both sites, is primarily of a marine source with varying contributions of terrestrial organic matter (inferred from the ratios of \( nC_{29}/nC_{30} \)).

4. Influxes of dinoflagellate, inferred from the high abundances of the 4-methylsteranes with the 23,24-dimethyl side chain, are significant at both sites.

5. The chemical organic matter in the Lower Cretaceous sediments from both sites is immature. This immaturity and the low levels of extractable hydrocarbons found in these sediments suggest caution when interpreting biomarker and degraded organic matter, which may not completely represent the total organic carbon content from the sediments.

6. Lower Cretaceous sediments from Argo are generally lean in organic matter, although a sample near the Barremian/Aptian boundary has a TOC of 5.1 wt% and samples near the Aptian/Albian boundary have TOC values up to 1.2 wt%. Sediments from the Aptian and upper Barremian contain marine components of organic matter and relatively high proportions of terrestrial organic matter.

7. Lower Cretaceous sediments from Gascouyse have low TOC (<0.1 wt%) in the Aptian/Albian and near the Barremian/Aptian boundaries. Maxima in TOC occur in the upper Barremian (1.02 wt%), the Barremian/Hauterivian (0.6 wt%), and the Valanginian (1.8 wt%). Sediments from Site 766 contain marine organic matter, and samples from the upper Barremian contain higher amounts of terrestrial organic carbon than older sediments.

8. Influxes of bacteria comprise significant contributions to the organic matter composition at both sites.

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