INTRODUCTION

Leg 75 of the Deep Sea Drilling Project (DSDP) was devoted to investigating the paleoenvironmental conditions prevailing in the South Atlantic ocean which have led to the accumulation of sediments rich in organic matter. One of the major objectives was to determine whether deposition occurred at the bottom of an isolated, anoxic basin or within a mid-depth oxygen-minimum zone. Previous deep sea drilling in the eastern South Atlantic during DSDP Leg 40 indicated the presence of numerous organic-matter-rich sediment sequences (Bolli, Ryan et al., 1978a,b). Cretaceous black shales were found in the Aptian–early Albian, in the late Albian, and in the Turonian–Santonian at DSDP Site 364 in the northern part of the Angola Basin and were interpreted as anoxic basin deposits (Bolli, Ryan et al., 1978b), whereas organic-matter-rich post–late Miocene sediments encountered in the Walvis Ridge area were suggested to be the result of the Benguela upwelling (Bolli, Ryan et al., 1978a).

During DSDP Leg 75, Hole 530A was drilled into the abyssal floor of the southeastern corner of the Angola Basin. The whole sediment sequence recovered was dominated by turbidites derived from the southwest African continental margin (Albian to Miocene) or from the Walvis Ridge (late Miocene to Pleistocene). Black shale layers in the Albian to Coniacian section were interbedded with red and green organic-matter-poor claystones.

Hydraulic piston cores from DSDP Hole 532, close to Site 362 on the Frio Ridge segment of the eastern part of the Walvis Ridge, represent an organic-matter-rich sequence of predominantly pelagic deposits with variable amounts of terrigenous clay. The organic matter in these late Miocene to Pleistocene sediments is supposed to be influenced by the Benguela upwelling system which started in the Miocene (see site summary, this volume).

RESULTS: HOLE 530A, ANGOLA BASIN

Although the seventeen core samples from Hole 530A represent most of the lithologic units, the investigation was concentrated mainly on lithologic Unit 8 (nine samples), which comprises the interbedded black-shale sequence of Albian–Cenomanian to Coniacian age. The organic carbon contents are moderate in the Miocene and younger sections (0.4–1.0%), very low in the upper Coniacian to Oligocene (<0.1%), and highly variable in Unit 8 (Table 1). Samples 530A-93-5, 135–145 cm and 530A-99-5, 0–10 cm contained thin black-shale layers, but these samples were processed without prior separation of the layers. Their organic carbon contents, thus, represent average values of different lithologies. In contrast to this, Sample 530A-97-3, 105–115 cm showed a clear boundary in the middle between a black shale in the top half and a green claystone. This sample was split, and subsamples of intervals 105–110 cm and 110–115 cm,
respectively, were analyzed separately. Organic carbon values of these subsamples show an extreme difference (i.e., 10.6% C$_{org}$ in the black shale and 0.10% C$_{org}$ in the green claystone).

The total extracts, normalized to organic carbon, reach values of 19 mg/g C$_{org}$, if the less accurate data of the organic-carbon-lean samples are disregarded. There is no clear downhole trend, which indicates that the organic matter in all samples is immature. There is, however, a slightly higher relative amount of total hydrocarbons (nonaromatic plus aromatic) in the more deeply buried Cretaceous samples than in Miocene and younger samples, indicating that diagenetic hydrocarbon formation is more advanced at greater depth.

Kerogen analysis by Rock-Eval pyrolysis (Espitalié et al., 1977) revealed that there are only a few samples with hydrogen index (HI) values above 100 mg HC/g C$_{org}$ (Fig. 1, Table 1), i.e., those samples from lithologic Unit 8 containing more than 2% organic carbon. The position of these samples in the Rock-Eval kerogen-type diagram (Fig. 1) corresponds to Type II kerogens, at a relatively high maturity stage. This is inconsistent with the low maturity of all the organic matter in the Hole 530A sediments (see discussion to come). For the kerogens involved, either higher hydrogen (less mature Type II) or higher oxygen index (OI) values (less mature mixed Type II/III) would have been expected. The reason for the actual results obtained is presently unknown. All the other samples from Hole 530A show low hydrogen index values and some very high oxygen index values, indicative of partly to strongly oxidized kerogen Type III organic matter. The Rock-Eval result of Sample 530A-34-6, 120-135 cm (HI: 272; OI: 405) seems to be erroneous or affected by contamination.

**Kerogen Microscopy**

**Maceral analysis.** The composition of the insoluble organic matter was studied by transmitted light, and normal and fluorescence reflected light microscopy of smear slides, polished separated kerogens and whole rock blocks. The results for eight samples from Hole 530A, six of which are from the Cretaceous interval, are shown in Figure 2. The other samples were very low in organic matter, so that the amounts of kerogen isolated were insufficient for microscopic investigation.

Most of the samples analyzed contain a considerable amount of degraded amorphous organic matter. Since, in the literature, different names have been attributed to the same type of unstructured liptinite, a brief definition will be given here (cf. Gormly and Mukhopadhyay, 1983). Bituminite II (corresponding to amorphinite B of van Gijzel, 1981; bituminite I of Teichmüller and Ottenjann, 1977; or bituminite of Robert, 1981) is a mostly grainy, dark gray amorphous mass of organic matter in normal reflected light, orange to dark reddish brown (at $R_m \approx 0.5\%$) in fluorescent light, and dark amber-yellow to light brown in transmitted light. This maceral type often contains very high amounts of framboïdal pyrite. The organic matter is assumed to be either the degradation product (as obvious from some relict structures) of phytosiga and zooplankton/benthos or of bacterially altered higher plant lipids.

Bituminite III (corresponding to amorphinite C of van Gijzel, 1981; and bituminite III of Teichmüller and Ottenjann, 1977) is also grainy, gray amorphous organic matter—sometimes with relict tissues of telinite or sporinite—in normal reflected light, showing higher reflectance than bituminite II, grayish brown to gray in fluorescent light (at $R_m \approx 0.5\%$), and dark gray to brown in transmitted light. This type of organic matter is probably the degradation product of humic matter and higher plant liptinites (mainly sporinite).

For a description of bituminite I (degraded algae), which we did not observe in the DSDP Leg 75 samples, we refer the reader to Gormly and Mukhopadhyay (1983).

The kerogen samples from the early Pliocene and late Miocene sections 530A-5-5 and 530A-9-2 are very similar to each other and show a predominance of humic and liptinite matter derived from terrestrial higher plants. Considerable amounts of huminite/vitrinite and bimacerite grains were found, with some of the vitrinites showing oxidation rims. Pollen and zooclasts are major constituents of both samples. The inertinites with high reflectance are mostly of the inertodetrinite type (Stach et al., 1975).

Among the Cretaceous samples investigated, those from Section 530A-93-5 (997.4 m, Coniacian) and Sec-

![Figure 1. Results of Rock-Eval pyrolysis, displayed as hydrogen index versus oxygen index diagram (Espitalié et al., 1977) for sediment samples from DSDP Holes 530A (Angola Basin, closed circles) and 532 (Walvis Ridge, open squares).](image-url)
tion 530A-99-5 (1050.1 m, Cenomanian) resemble each other, both in transmitted and reflected light. They contain neutral gray-black amorphous humic and liptinite material as dominant macerals (bituminite III and II), often enriched with organo-pyrite, which in some cases show pseudomorphism on plant tracheids, pollen, and bryozoan fragments. Another major component in both samples is sporinite from pollen and spores (Plate 1, Fig. 3). The Cenomanian kerogen (530A-99-5) is enriched in both autochthonous (0.25–0.65% reflectance) and recycled vitrinite (0.75–1.55% reflectance), the latter showing oxidation rims and bimaculate grains. In contrast to
this, very few recycled vitrinite particles were observed in Section 53OA-93-5 (Coniacian). The overall composition of both samples indicates a dominant contribution of terrigenous organic matter mixed with a minor component of marine origin.

The Cenomanian black shale (Sample 530A-97-3, 105-110 cm) contains more than 80% mixed amorphous organic matter. In transmitted light, there is a mixture of amber yellow and gray-black amorphous organic matter (Plate 1, Figs. 1, 4, 5), with some slightly degraded and some intact dinoflagellates (Plate 1, Figs. 1, 5). In reflected light, dark gray bituminite II (Plate 1, Fig. 2) dominates, bearing a high content of organo-pyrite pseudomorphs (observed in whole rock specimen) on zoo- and phytoplankton and occasionally on pollen grains. Fluorescent reflected light predominantly shows reddish brown bituminite II. In summary, this kerogen is of mixed marine and terrigenous origin with phyto- and zooplankton dominating. Colonial or unicellular algae with grain boundaries are absent.

The Cenomanian green claystone, Sample 530A-97-3, 110-115 cm, which is from the lower contact of the black shale discussed above contains a mainly fusinized amorphous organic matter with subordinate organo-pyrite. Dominant macerals in normal reflected light are granular micrinite and degradoinertinite (Stach et al., 1975).

The late Albian-early Cenomanian gray shale from Section 53OA-103-3 (1084.3 m) has a kerogen similar in appearance to that of the Cenomanian black shale. Gray bituminite III (Plate 1, Fig. 6) and bituminite II (Plate 1, Fig. 7) with a very high content of frambooidal pyrite dominate in normal reflected light. A few liptinite particles are seen in addition to the bulk dark gray amorphous organic matter in transmitted light.

The purple gray shale of late Albian-early Cenomanian age (Section 53OA-105-1) contains a mixture of bituminite III, bituminite II, sporinite, and phyto- and zooclasts (Plate 1, Figs. 7, 8). The dominant source for this organic matter is terrestrial higher plants.

In Figure 3 the results of the maceral analysis are correlated with the hydrogen index values of the Rock-Eval pyrolysis. Three groups of organic-matter types can be distinguished. The Cenomanian black shale is richest in marine macerals and hydrogen, although for a Type II kerogen (cf. Tissot and Welte, 1978), with the given ratio of marine versus terrigenous organic matter, the hydrogen index value appears to be unusually low. The sample from Section 530A-103-3 contains a mixed kerogen of Type II-III, whereas all other samples from Hole 530A contain a dominance of terrigenous organic matter which is partly oxidized, resulting in low hydrogen index values. Only the Type II-III and Type II kerogen samples bear a significant petroleum generation potential.

Maturity. Figure 4 shows reflectance histograms of autochthonous (primary) and recycled vitrinite/huminite for sediment samples from DSDP Hole 530A (Angola Basin).
Nonaromatic Hydrocarbon Composition

With the organic-carbon-rich black shale sample (Sample 530A-97-3, 105–110 cm) as the only exception, all nonaromatic hydrocarbon distributions are dominated by n-alkanes (Fig. 5 A–H; Table 2). Within the C_{15}+ n-alkanes, the long-chain members with 29, 31, and 33 carbon atoms predominate in all samples. The maximum is at n-C_{31} in each case, and there is only a slight decrease in even-over-odd carbon-number predominance down the hole (cf. Figs. 5A and 5B).

A series of 17β(H)-hopanes (C_{27}, C_{28}, C_{32}; maximum at C_{31}) are the only cyclic hydrocarbons present in significant, albeit low, concentrations in the moderately organic-carbon-rich Miocene and younger sediments from Hole 530A (Fig. 5A). In addition, traces of C_{32} ster-4-enes and ster-5-enes, hop-17(21)-ene, fern-8-ene, fern-9(ll)-ene, and 17α(H)-hopanes (C_{27}, C_{29}, C_{30}) could be detected by gas chromatography/mass spectrometry (see chapter appendix for compound structures). The organic-carbon-lean sediments, represented by Samples 530A-87-3, 120–130 cm (Coniacian; Fig. 5B) and 530A-97-3, 110–115 cm (Cenomanian; Fig. 5E), are nearly devoid of any nonaromatic hydrocarbons, except n-alkanes and some pristane and phytane.

The nonaromatic hydrocarbon compositions of those sediment samples from lithologic Unit 8 containing between 0.7 and 2.8% C_{org}, i.e., Samples 530A-93-5, 135–145 cm (Coniacian; Fig. 5C); 530A-99-5, 0–10 cm (Coniacian; Fig. 5F); 530A-103-3, 110–120 cm (Albian–Cenomanian; Fig. 5G); and 530A-105-1, 90–100 cm (Albian–Cenomanian; Fig. 5H), are similar to each other with respect to the type of compounds present, although there are distinct variations in relative concentrations. The content of cyclic compounds relative to n-alkanes does not always follow the total organic carbon trend; e.g., the highest relative concentration of cyclic nonaromatic hydrocarbons is found in Sample 530A-105-1, 90–100 cm (Fig. 5H) with only 0.82 C_{org}. Rearranged sterenes (C_{27}–C_{29}) are the only steroid hydrocarbons present, and these are less abundant than triterpenoid hydrocarbons. 17β(H)-hopanes (C_{27}, C_{28}, C_{32}) are still abundant in the deepest sample from 1095.0 m (Fig. 5H) which, however, also contains 17α(H)-trisnorhopane in unusually high concentration. Triterpenes are common in all the samples discussed here and compose a series of hop-17(21)-enes (C_{27}, C_{29}, C_{30}), hop-13(18)-enes (C_{27}, C_{29}, C_{30}), fern-8-ene, fern-9(ll)-ene, and a Δ^{15}-triterpene. An unknown major compound (a' in Fig. 5 may also be a C_{30}-triterpene (molecular ion at m/z 410, base peak m/z 395).

The Cenomanian black shale, Sample 530A-97-3, 105–110 cm, differs from the other samples from Unit 8 in the fact that phytane and pristane are the dominant compounds in the nonaromatic hydrocarbon fraction, and sterols and triterpenoids are abundant in the higher molecular weight range (Fig. 5D). The distribution of triterpenes and triterpanes present are qualitatively similar to the other samples from lithologic Unit 8; but with the exception of the unknown compound a', are less abundant than the steroid hydrocarbons; among the latter, the rearranged steranes have the highest concentration (Fig. 6). In addition 4-methyl rearranged steranes (m/z 271) and diunsaturated compounds (m/z 269) add to the complex sterane pattern (Fig. 6). C_{27} and C_{29} ster-4-enes and ster-5-enes (m/z 215) are minor constituents, as are the saturated C_{27} to C_{29} 5α(H)- and 5β(H)-steranes (m/z 217; Fig. 6).

Aromatic Hydrocarbon Composition

Six aromatic hydrocarbon fractions of sediments from Hole 530A were analyzed in detail; the results are shown in Figures 7 A–F and Table 3. The late Miocene Sample 530A-9-2, 120–130 cm contains different compounds from those present in samples from lithologic Unit 8; three different alkylated thiophenes are major constituents. The mass spectra of two isomeric C_{20} thiophenes are shown in Figure 8; they are consistent with the general fragmentation pattern of alkylated thiophenes (Kinney and Cook, 1952; Pomonis et al., 1976). A tentative structure of 2-pentadecanoyl-3-methylthiophene was assigned for one isomer, where the pentadecanoyl substituent may be a straight chain or branched, although not at the benzylic position. The second thiophene is mono-substituted. The formation of the main fragment at m/z 98 is supposed to result from a McLafferty rearrangement common in long-chain alkylated aromatics unless both ortho-positions are devoid of hydrogens, as in the case of the first isomer (Budzikiewicz et al., 1967), although in the mass spectra of 2-n-tridecanyl- and 2-n-heptadecanoyl-thiophene published by Pomonis et al. (1976) the m/z 98 fragment shows only half the abundance of the m/z 97 fragment. The extent of rearrangement vs. cleavage products may, however, depend on instrumental parameters. The mass spectrum of the third alkylated thio phosphine (C_{27}; 49 in Fig. 7A) has been reported previously (Rulkotter, von der Dick, and Welte,

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Table 2. Nonaromatic hydrocarbons identified by their mass spectra and relative retention times in sediments from DSDP Holes 530A and 532 (cf. Figs. 5 and 12).

| p' = pristane | w = 24-methylcholesta-4,22-diene |
| q = C_{29} diasterene | x = 24-methylcholesta-5,22-diene |
| r = 17β(H)-bishomohopane | y = 17α(H)-trisnorhopane |
| s = 17/3(H)-bishomohopane | z = 17β(H)-22,29,30-trisnorhopane |
| t = 17j3(H)-homohopane | a' = unknown (M+410, BP 395) |
| u = cholest-5-ene | b' = 30-norhop-13(18)-ene |
| v = unknown (M+410, BP 395) | c' = 24-ethylcholest-4-ene |
| b = C_{25} isoprenoid | d' = 17α(H)-30-norhopane |
| c = unknown C_{27} sterene | e' = 24-ethylcholest-5-ene |
| d = C_{27} diasterene | f' = 30-norhop-13(18)-ene |
| e = unknown C_{29} sterene | g' = 30-norhop-13(18)-ene |
| f = C_{27} diasterene | h' = hop-17,21(1)-ene |
| g = C_{27} diasterene | i' = 30-normoretane |
| h = C_{27} diasterene | j' = 17α(H)-hopane |
| i = unknown C_{27} sterene | k' = fern-8-ene |
| j = unknown (M+398, BP 231) | l' = hop-17,21(1)-ene |
| k = squalanet(?) | m' = 30-norhop-13(18)-ene |
| l = C_{28} diasterene | n' = 30-normoretane |
| m = C_{28} diasterene | o' = C_{30} Δ^{15}-triterpene |
| n = C_{28} diasterene | p' = fern-8-ene |
| o = unknown C_{29} sterene | q' = C_{30} Δ^{15}-triterpene |
| p = unknown (BP 231) | r' = 17β(H)-trisnorhopane |
| q = C_{29} diasterene | s' = C_{30} Δ^{15}-triterpene |
| r = C_{29} diasterene | t' = 17β(H)-trisnorhopane |
| s = cholest-4-ene | u' = cholest-5-ene |
| t = C_{29} diasterene | v' = C_{30} Δ^{15}-triterpene |
| u = cholest-5-ene | w' = C_{30} Δ^{15}-triterpene |
| v = 5α(H)-cholestane | x' = C_{30} Δ^{15}-triterpene |

Note: BP = base peak
Figure 5. Capillary column gas chromatograms of the nonaromatic hydrocarbon fractions of selected sediment samples from DSDP Hole 530A (Angola Basin). Numbers indicate n-alkanes, compounds marked with letters are listed in Table 2, pr = pristane, ph = phytane, x = phthalate ester contaminant. Analytical conditions: 25 m fused silica capillary column coated with SE 54; temperature program: 100°-300°C at 3°C/min.; slightly different conditions for chromatogram in Figure 5D (wider bore glass capillary column, temperature program 80-250°C). (Revisions of stratigraphic ages are given in Steinmetz et al., this volume.)
The late Miocene sample is the only one from Hole 530A containing significant amounts of perylene. Two isomeric tetramethyloctahydrochrysenes probably derived from oxidation products of pentacyclic triterpenoids which have lost their A-ring (Spyckerelle et al., 1977a,b). Two further aromatic hydrocarbons have molecular weights of 326 and 310, respectively. They could not be identified with certainty, although the latter may be a monoaromatic pentamethyldodecahydrochrysenes tentatively identified in Messel oil shale by Spyckerelle et al. (1977a).

The Cretaceous sediments from lithologic Unit 8 have aromatic hydrocarbon compositions similar to each other, with the exception of the black shale, Sample 530A-97-3, 105–110 cm (Figs. 7B–F). The black shale shows a complex, poorly resolved aromatic hydrocarbon mixture in the lower molecular weight range (Fig. 7C), with only one major peak which gives a mass spectrum virtually identical with that of 1,1,6-trimethyl-1,2,3,4-tetrahydronaphthalene (ionene). In the higher molecular weight range, monoaromatic steroid hydrocarbons dominate. In addition to some unknown com-

Figure 5. (Continued).
represented by the m/z 231 trace, and an additional
and Welte, 1983). Triaromatic steroid hydrocarbons are
211 fragmentogram may be anthrasteroids (Rullkötter
whereas the two earlier eluting compounds in the m/z
27
concentration as C
homologs. 4-Methylstera-
29
are less stable than the C-ring aromatic analogs (Rull-
and ring B monoaromatic steroid hydrocarbons which
The aromatic steroid hydrocarbon distribution of Sample
37-2 having a lower hydrogen index value than expected
37-2 of the Rock-Eval pyrolysis classified all kerogens as mixed marine/terrigenous (Type II–III), with only the sample from Section 532-
3 having an m/z 245 key fragment is also present (not shown in Fig. 9).
RESULTS: HOLE 532, WALVIS RIDGE
The total organic carbon content of Pliocene to Pleis-
tocene sediments from DSDP Hole 532 are consistently high, with a variation between 1.92 and 4.40% (Table 3). Rock-Eval pyrolysis indicates a mixed Type II/Type III (planktonic/terrestrial higher plant) kerogen with little variation for all samples, except the deepest of early Pli-
cene age, which has a lower hydrogen index and would be classified as Type III kerogen (Fig. 1). The total extracts, normalized to organic carbon, show a clear differ-
Kerogen Microscopy
Maceral analysis. The maceral composition of the
Hole 532 samples, derived from transmitted and reflect-
ed light microscopy, is shown in Figure 10. Petrographic
data indicate that all six kerogens are similar in appear-
ance and composition. More than 75% of the organic
matter in each case is composed of degraded, amorphous
bituminite II and III, with bituminite II being more than
50%, except in Section 532-13-2. Plate 1, Figure 12
shows the bituminite II in Sample 432-37-2, 110-120 cm; this sediment is richest in this type of organic matter. In
general, all kerogens show a grunny gray mass of organic
matter with a high concentration of frambooidal pyrite.
Some samples contained relict degraded dinoflagellates.
The content of huminite/vitrinite and inertinite particles
is low throughout. The maceral composition plotted
against the hydrogen index of the Rock-Eval pyrolysis
classifies all kerogens as mixed marine/terrigenous
(Type II–III), with only the sample from Section 532-
37-2 having a lower hydrogen index value than expected
from the maceral composition (Fig. 4).
Maturity. The reflectance histograms of autochthon-
ous (primary) and recycled huminite/vitrinite (Fig. 11)
exhibit multimodal distributions, as in the Angola Basin
samples from DSDP Hole 530A. For the fragmentary
vitrinites in Section 532-20-2, reliable reflectance mea-
surements could not be achieved. The mean huminite/
vitrinite reflectance values show a slight increase with
depth (Fig. 10), but indicate that the organic matter is
immature in all samples.
Nonaromatic Hydrocarbon Composition
The n-alkane distributions of all samples from Hole
532 exhibit major similarities. Terrigenous influence is
demonstrated by the dominance of long chain-length
wax alkanes (Eglinton and Hamilton, 1963), with a max-
imum at n-C
and a high odd-over-even carbon number predominance (Fig. 12). Isoprenoid hydrocarbons are
represented by small amounts of pristane, phytane, a
C
-isoprenoid and, in one sample, probably squalane.
In addition, the more deeply buried samples contain
some phytanes (Figs. 12C and D). Cyclic hydrocarbons
are also minor constituents and predominantly consist
of methylated series having an m/z 245 key fragment is al-
so present (not shown in Fig. 9).

Figure 6. Mass fragmentograms of key fragment ions of steroid hy-
drocarbons in the nonaromatic hydrocarbon fraction of the black
shale Sample 530A-97-3, 105-110 cm (see text). Compounds marked
with letters are listed in Table 2. (Revisions of stratigraphic ages
are given in Steinmetz et al., this volume.)
of 17β(H)-hopanes and triterpenes. Sterenes and steradienes are present in the two deepest samples, whereas diasterenes could not be detected. The early Pliocene sample, Sample 532-37-2, 110–120 cm, contains a series of unknown isomeric compounds with m/z 231 as the only major ion in their mass spectra.

### Aromatic Hydrocarbon Composition

Three samples from Hole 532 were analyzed for their aromatic hydrocarbon distributions, and there is a clear difference between the shallowest Pleistocene sample (Sample 531-1-2, 110–120 cm; 2.7 m) and the Pliocene sediments (Fig. 13). The former apparently contains a number of nitrogen-bearing aromatics as major components (19, 30–32) according to their odd-mass molecular ion peaks (Fig. 13A; Table 4). Alkyl thiophenes (20, 21, 40, 49) are common in all samples, although in Sample 532-1-2, 110–120 cm only the C\textsubscript{20}-compound (40) could be detected. A number of steroid hydrocarbons, with various degrees of unsaturation, are dominant in the higher molecular weight region of the Pliocene samples (Figs. 13B and C). Some of them have mass spectra resembling sterenes and might have been expected to occur in the nonaromatic hydrocarbon fraction. Perylene (43) is present in samples from the Pliocene in significant concentration.

### DISCUSSION: HOLE 530A, ANGOLA BASIN

#### Organic Facies

The results of the organic geochemical and organic petrographic investigation of the DSDP Leg 75 core samples from Hole 530A in the Angola Basin indicate that the environmental conditions were highly variable with respect to organic matter preservation over the time span covered, i.e., from the middle Cretaceous to the late Tertiary. The Coniacian (Core 530A-89) through Oligocene sediments are poor in organic matter and represent typical deep-sea sediments deposited in oxidizing water (Table 1). The residual organic matter present consists of highly oxidized, inert material predominantly of terrigenous origin, as shown by Rock-Eval pyrolysis (Fig. 1; Table 1; Welte et al., 1979). The nonaromatic hydrocarbon fractions (e.g., Fig. 5B) contain mostly n-alkanes, with a distribution typical of higher plant waxes (Eglinton and Hamilton, 1963). These compounds are known to survive long distal transport into the deep-sea area (e.g., Rullkötter et al., 1980; Rullkötter, Cornford, Welte, 1982).

The Miocene and Pliocene sediments have organic carbon contents between 0.4 and 1.0% (Table 1), and contain organic matter of predominantly terrigenous origin which is slightly better preserved than in the Coniacian through Oligocene sediments previously described. Kerogen microscopy reveals the presence of only 20–30% marine organic matter (Fig. 2), and Rock-Eval pyrolysis (Fig. 1, Table 1) supports this by yielding low hydrogen index values and high oxygen index values around 300 mg CO\textsubscript{2}/g C\textsubscript{org}. The nonaromatic hydrocarbon fractions (e.g., Figure 5A) are dominated by n-alkanes derived from terrestrial higher plants. Small amounts of cyclic compounds are mainly triterpanes of the 17β(H)-hopane type and indicate partial microbial reworking of the organic matter at the time of deposition (Ourisson et al., 1979). The late Miocene sample, 530A-9-2, 120–137 cm (Fig. 7A) contains, in addition to the thiophenes of unknown significance, two degraded triterpanes of the octahydrochrysene type as major compounds of the aromatic hydrocarbon fraction. These are thought to be derived from oxidative degradation of 3-oxo triterpenoids (Corbet et al., 1980) common in terrestrial higher plants. This further confirms that the African continent was the main source of organic matter in these sediments, and that the organic matter underwent partial oxidation during transport. The Benguela upwelling current, which started in the Miocene, apparently has not had a pronounced effect on the Miocene/Pliocene sediments at Site 530 in the Angola Basin. This means that the oxygen minimum zone usually associated with high bioproductivity in areas of coastal upwelling did not extend far enough, either laterally or vertically, to cause a significant preservation of the autochthonous marine biomass at this site. Compared with the organic-carbon-lean Coniacian through Oligocene sediments at this site, the better preservation of the organic matter may be the result of higher sedimentation rates and rapid burial into turbidity deposits, probably originating from the nearby Walvis Ridge (cf. site summary, this volume).

The middle Cretaceous sequence with alternating organic-carbon-rich black shales and organic-carbon-lean claystones illustrates two extreme modes of deposition and preservation of organic matter in the deep sea. The organic-carbon-lean claystones, exemplified by the Cenomanian sample, Sample 530A-97-3, 110–115 cm, bear the same characteristics as the Coniacian through Oligocene sediments described before, which all contain residual amounts of organic matter below 0.1%. The Cenoo-

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**Table 3. Lithology, stratigraphy, organic carbon content, extract and liquid chromatography yields, and Rock-Eval pyrolysis data.**

<table>
<thead>
<tr>
<th>Sample (interval in cm)</th>
<th>Depth (m)</th>
<th>Lithology</th>
<th>Stratigraphic age</th>
<th>C\textsubscript{org} (%)</th>
<th>Extract (mg/g C\textsubscript{org})</th>
<th>Chromatographic fractions (%)</th>
<th>Rock-Eval pyrolysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-2, 110-120</td>
<td>2.7</td>
<td>Nannofossil-foraminiferal ooze</td>
<td>late Pleistocene</td>
<td>2.76</td>
<td>4</td>
<td>Nonaromatic 19, Aromatic 6.1, Het-comp. 3.4, Residue 1.3</td>
<td>HI 2.73, OI 26, Tmax 394</td>
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<tr>
<td>7-9, 90-100</td>
<td>29.5</td>
<td>Foraminiferal-nannofossil ooze</td>
<td>early Pleistocene</td>
<td>4.40</td>
<td>1.2</td>
<td>Nonaromatic 3.0, Aromatic 6.1, Het-comp. 4.0, Residue 5.4</td>
<td>HI 361, OI 57, Tmax 359</td>
</tr>
<tr>
<td>13-2, 90-100</td>
<td>54.9</td>
<td>Diatomaceous-nannofossil marl</td>
<td>early Pleistocene</td>
<td>3.07</td>
<td>3.9</td>
<td>Nonaromatic 2.0, Aromatic 4.1, Het-comp. 2.0, Residue 2.7</td>
<td>HI 208, OI 20, Tmax 317</td>
</tr>
<tr>
<td>20-2, 120-130</td>
<td>86.0</td>
<td>Nannofossil-diatomaceous marl</td>
<td>late Pleistocene</td>
<td>3.59</td>
<td>2.0</td>
<td>Nonaromatic 0.4, Aromatic 3.0, Het-comp. 0.2, Residue 0.5</td>
<td>HI 286, OI 28, Tmax 352</td>
</tr>
<tr>
<td>29-2, 90-100</td>
<td>123.3</td>
<td>Nannofossil marl</td>
<td>late Pliocene</td>
<td>3.92</td>
<td>2.0</td>
<td>Nonaromatic 0.4, Aromatic 3.0, Het-comp. 0.2, Residue 0.5</td>
<td>HI 286, OI 28, Tmax 352</td>
</tr>
<tr>
<td>37-2, 110-120</td>
<td>193.3</td>
<td>Foraminiferal-nannofossil ooze</td>
<td>early Pliocene</td>
<td>1.92</td>
<td>1.9</td>
<td>Nonaromatic 0.4, Aromatic 3.0, Het-comp. 0.2, Residue 0.5</td>
<td>HI 109, OI 10, Tmax 352</td>
</tr>
</tbody>
</table>
Figure 7. Capillary column gas chromatograms of the aromatic hydrocarbon fractions of selected sediment samples from DSDP Hole 530A (Angola Basin). Numbered compounds are listed in Table 3; x = phthalate ester contaminant. Analytical conditions as for Figure 5 (starting and end temperature slightly variable). (Revisions of stratigraphic ages are given in Steinmetz et al., this volume.)
manian green claystone contains more than 60% inerti-
nite particles (Fig. 2) and only a minor marine organic
matter fraction. The nonaromatic hydrocarbons are
dominated by higher plant wax n-alkanes (Fig. 5E).
These data suggest that deposition occurred under oxi-
dizing deep-water conditions as indicated by the deposi-
tion of the red claystones as autochthonous pelagic and
partly turbiditic background sediment (cf. site sum-
mary, this volume).

The overlying black shale, with an organic carbon
content one hundred times higher than the green clay-
stone according to shipboard sedimentological observa-
tions (site summary, this volume), represents a turbidite
deposit. It derived from a zone of oxygen-depleted wa-
ter where primary deposition occurred under conditions
favorable for the preservation of large amounts of or-
ganic matter. Flow of debris down slope on steep con-
tinental margins, combined with an oxygen-minimum
zone in the outer shelf or upper slope area as a result of
high bioproductivity, has already been shown to be an
important mechanism for the deposition of sediments,
rich in organic matter, in the deep sea (Demaison and
Moore, 1980; Rullkötter et al., 1983). In the case of
the Cenomanian black shale, nearly 80% of the organic
matter appears to be of marine origin. Most of it ap-
ppears to be affected by microbial reworking, and is thus
detected as amorphous bituminite II (Fig. 2). The ter-
rigenous influence, however, is still clearly visible both
by kerogen microscopy (Fig. 2) and the n-alkane distri-
bution in the nonaromatic hydrocarbon fraction (Fig.
5D) which, as in all other Hole 530A sediments, has a
maximum at n-C\textsubscript{31} and a strong even-over-odd carbon
number predominance. The relatively low hydrogen in-
dex from Rock-Eval pyrolysis (Fig. 3) may be an effect
of the strong microbial reworking. The high pristane
and phytane concentrations in the black shale sample
(Fig. 5D) can also be taken as an indication of a deposi-
tional environment favorable for the preservation of
(marine) organic matter (cf. composition of deep sea
sediments from the outer California borderland, Rull-
kötter et al., 1981; and of a black shale from DSDP Site
367, Rullkötter, Cornford, and Welte, 1982). The domi-
nance of rearranged steroid hydrocarbons, indicating
high catalytic activity of the clay matrix (Sieskind et al.,
1979), is common in black shales and has also been
found in black shales from the North Atlantic (Rullköt-
ter, Cornford, and Welte, 1982) and the Falkland Pla-
teau in the Southeast Atlantic (von der Dick et al.,
1983).

The remainder of the Albian/Cenomanian to Conia-
cian (Core 530A-93) sediments exhibit an intermediate
organic facies between the black shale and the green
claystone from Section 530A-97-3, some of them even
representing sediments of these extremes that could not
be separated. The content of macerals of marine origin
roughly follows the total amount of organic carbon

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Figure 7. (Continued).
J. RULLKÖTTER, P. K. MUKHOPADHYAY, D. H. WELTE

Figure 8. Mass spectra of two isometric C20 alkyl thiophenes. Structures are tentatively derived from mass spectrometric fragmentation only.

present but, except in Section 530A-103-3, terrigenous components dominate (Fig. 2). This indicates an effect of selective mineralization of lipid components, where marine lipids are more easily destroyed or altered than terrigenous lipids. An explanation for this may be better protection of higher plant lipids by their specific cell wall composition or by structural components which are not readily degradable. All nonaromatic hydrocarbon fractions show n-alkanes as the most abundant compounds. In contradiction to the microscopic results, Sample 530A-105-1, 90–100 cm has the highest relative concentration of branched and cyclic nonaromatic hydrocarbons rather than Sample 530A-103-3, 110–120 cm (Fig. 5 G, H). In the aromatic hydrocarbon fractions of all these sediments, degraded polycyclic terpenoids are abundant components. Among them are nordehydroabietane or an isomer (peak No. 16 in Fig. 7B, E, F), a monoaromatic tetracyclic compound (peak No. 27), and the pentacyclic 7-methyl-3'-ethyl-1, 2-cyclopentanochrysene (peak No. 54). The mass spectrum of the tetracyclic compound fits with the hypothetical structure (27) tentatively assigned to a constituent of the Messel oil shale by Spyckerelle et al. (1977a, b). The cyclopentanochrysene (54) probably is related to the hopane triterpenoids found in the nonaromatic hydrocarbon fractions and indicates bacterial activity (Ourisson et al., 1979).

Diagenesis

The huminite/vitrinite reflectance measurements show a maturity increase from about 0.3% Rm in the Tertiary samples around 200 m depth to 0.48% Rm in the Albian–Cenomanian at 1095.0 m depth. No trend could be established because of the large gap between 200 m and nearly 1000 m depth, where measurements were prevented by the low organic carbon concentrations. The same argument affects any detailed discussion of diagenetic alterations in the extractable hydrocarbon fractions. There is a decrease in the odd-over-even carbon number preference (calculated as 2 C31/C30 + C32) from the late Miocene sample at 203.8 m (11.1 for the C31 n-alkane) to the Cretaceous sediments, where the CPI-values range between 6 and 8 for most of the samples, except Sample 530A-97-3, 110–115 cm which has a CPI-value of 3.7, possibly affected by the high content of reworked organic matter (Fig. 5). The detailed hydrocarbon distributions within the series of Cretaceous samples are apparently more influenced by differences in source organic matter composition than by temperature-related diagenetic alterations over a depth interval of less than 100 m.
ORGANIC MATTER IN SEDIMENTS

Organic facies (percent of maceral type)

Sub-bottom depth (m)

Vitrinite reflectance

Huminite/vitrinite
Bituminite III
Inertinite
Particulate (higher plant) lipids
Phyto- and zooclasts other than algae
Bituminite II

Figure 10. Maceral composition and mean vitrinite reflectance for sediment samples from DSDP Hole 532 (Walvis Ridge).

DISCUSSION: HOLE 532, WALVIS RIDGE

The six sediment samples of Pliocene to Pleistocene age from the Walvis Ridge are fairly uniform in maceral (Fig. 10) and nonaromatic hydrocarbon composition (Fig. 12), whereas in the aromatic hydrocarbon fractions analyzed (Fig. 13), the shallowest sample (2.7 m; late Pleistocene) differs from the other two, probably because of a diagenetic effect. The relatively high organic carbon content in all samples is consistent with a high percentage of marine organic matter, although it is mostly present as amorphous (degraded) bituminite II (Fig. 10). The high amount of organic matter preserved may be related to high primary bioproductivity within the Benguela upwelling regime, but may also be a result of the high sedimentation rates (cf. site summary, this volume). Transport of terrigenous organic matter from the nearby African continent, nonetheless, has been significant over the whole time span. This is indicated both by the maceral analysis (Fig. 10) and the gas chromatograms of the nonaromatic hydrocarbon fractions (Fig. 12) where, partly as a result of the low maturation level, higher plant wax alkanes clearly dominate. The mixed organic matter composition is further supported by the intermediate hydrogen and oxygen index values (Figs. 1 and 3) obtained from the Rock-Eval pyrolysis.
Figure 12. Capillary column gas chromatograms of the nonaromatic hydrocarbon fractions of selected sediment samples from DSDP Hole 532 (Walvis Ridge). Numbers denote n-alkanes, lettered compounds are listed in Table 2; pr = pristane, ph = phytane, x = phthalate ester contaminant. Analytical conditions as for Figure 5.
ORGANIC MATTER IN SEDIMENTS

Figure 13. Capillary column gas chromatograms of the aromatic hydrocarbon fractions of selected sediment samples from DSDP Hole 532 (Walvis Ridge). Numbered compounds are listed in Table 3. Analytical conditions as for Figure 5.

CONCLUSIONS

All sediments from the Angola Basin, DSDP Site 530, and the Walvis Ridge, DSDP Site 532, are strongly influenced by the seaward transport of terrigenous organic matter from the African continent. Under typical deep sea conditions in oxidizing waters, only a small fraction of this material survives, and most is strongly oxidized during transport; virtually all the marine organic matter is destroyed, leading to sediments with usually less than 0.1% organic carbon. Only down-slope mass transport with, first, high sedimentation and high burial rates and, second, a high content of organic matter in the sediment material from primary deposition in an oxygen-depleted zone leads to deep sea sediments enriched in organic carbon at DSDP Site 530 in the Angola Basin. The amount of marine organic matter preserved usually follows the total organic carbon concentrations. In extreme cases, black shales with more than 10% organic carbon and a hydrogen-rich Type II kerogen are deposited. At Site 532, Walvis Ridge, the high organic carbon contents, which represent a mixture of terrigenous and marine organic matter, are thought to be related to the high bioproductivity in the Benguela upwelling regime and the high sedimentation rates. Organic matter in all organic-carbon-rich samples investigated seems to be strongly reworked by microbial action as indicated by the dominance of bituminite II and III. The maturation level of the organic matter is low throughout.

ACKNOWLEDGMENTS

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REFERENCES


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Table 4. Aromatic hydrocarbons, identified or characterized by major peaks in their mass spectra, occurring in sediments from DSDP Holes 530A and 532 (cf. Figs. 7 and 13).

<table>
<thead>
<tr>
<th>No.</th>
<th>Compound</th>
<th>M⁺</th>
<th>Major mass spectral peaks (rel. intensity, %)</th>
</tr>
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<tr>
<td>1</td>
<td>Methylbenzaldehyde</td>
<td>142</td>
<td>—</td>
</tr>
<tr>
<td>2</td>
<td>Cyclohexadienone (constituent?)</td>
<td>166</td>
<td>166(15), 83(50), 82(100)</td>
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<tr>
<td>3</td>
<td>Iones (1,1,4,5-tetramethyl-1,2,3,4-tetracyclicalphane)</td>
<td>174</td>
<td>174(18), 159(100)</td>
</tr>
<tr>
<td>4</td>
<td>C₆-naphthalene</td>
<td>156</td>
<td>—</td>
</tr>
<tr>
<td>5</td>
<td>C₇-naphthalene</td>
<td>170</td>
<td>—</td>
</tr>
<tr>
<td>6</td>
<td>2-methylindanyl</td>
<td>168</td>
<td>168(100), 167(50), 165(50), 153(60)</td>
</tr>
<tr>
<td>7</td>
<td>C₈-naphthalene</td>
<td>184</td>
<td>—</td>
</tr>
<tr>
<td>8</td>
<td>Phenanthrene</td>
<td>178</td>
<td>—</td>
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<td>9</td>
<td>Alkylbenzene</td>
<td>246</td>
<td>246(15), 106(100), 105(50)</td>
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<td>246</td>
<td>246(10), 120(50), 119(100)</td>
</tr>
<tr>
<td>11</td>
<td>Alkylbenzene</td>
<td>376(7)</td>
<td>119(100)</td>
</tr>
<tr>
<td>12</td>
<td>Methylphenanthrene</td>
<td>192</td>
<td>—</td>
</tr>
<tr>
<td>13</td>
<td>Unknown</td>
<td>256</td>
<td>256(20), 241(100)</td>
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<td>Unknown</td>
<td>208</td>
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<td>Unknown</td>
<td>208</td>
<td>208(100), 207(52), 193(10), 179(14), 178(13), 165(29)</td>
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<td>16</td>
<td>Norbornadiene?</td>
<td>256</td>
<td>256(35), 241(75), 171(63), 159(68), 145(100), 69(34)</td>
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<td>17</td>
<td>Pyrene</td>
<td>202</td>
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<td>Alkylbenzene</td>
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<td>106(100), 105(60)</td>
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<td>19</td>
<td>Unknown</td>
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<td>20</td>
<td>Alkylhydrocarbon</td>
<td>308</td>
<td>cf. Fig. 8a</td>
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<td>Alkylbenzene</td>
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<td>316(15), 106(100), 105(70)</td>
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<td>Unknown</td>
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<td>326(5), 311(30), 106(100), 95(90)</td>
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<td>26</td>
<td>Pentamethyldodecahydrocrysene?</td>
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<td>27</td>
<td>3,3',7,12a-tetraethyl-1,2,3,4,4a,11,12,12a-octahydrocrysene</td>
<td>292</td>
<td>292(100), 207(20), 192(15), 181(12), 168(17)</td>
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<tr>
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<td>Isomer of 29 (3,4,7,12a-tetramethyl-1,2,3,4,4a,11,12,12a-octahydrocrysene)</td>
<td>292</td>
<td>292(100), 207(47), 207(95), 193(15), 186(30)</td>
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<td>Unknown</td>
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<td>207(86), 206(22), 129(62), 105(18), 91(100)</td>
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<td>31</td>
<td>Unknown</td>
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<td>207(95), 206(18), 129(58), 105(15), 91(100)</td>
</tr>
<tr>
<td>32</td>
<td>C-ring aromatic steroid (BP 239)</td>
<td>—</td>
<td>239(100)</td>
</tr>
<tr>
<td>33</td>
<td>C-ring aromatic steroid (BP 253)</td>
<td>—</td>
<td>253(100)</td>
</tr>
<tr>
<td>34</td>
<td>3,3',7-trimethyl-1,2,3,4-tetrahydroxylene</td>
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<td>274(60), 218(100)</td>
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<td>Unknown</td>
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<td>400(20), 175(23), 135(100)</td>
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<td>36</td>
<td>Unknown</td>
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<td>410(44), 397(75), 377(16), 176(15), 149(27), 123(35), 109(42), 81(100)</td>
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<td>Steradiene? (C₁₇)</td>
<td>368</td>
<td>368(100)</td>
</tr>
<tr>
<td>42</td>
<td>Perylene</td>
<td>252</td>
<td>252(100)</td>
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<tr>
<td>43</td>
<td>Sterrene? (C₂₃)</td>
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<td>380(80), 265(65), 105(63), 81(95), 69(95), 55(100)</td>
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<td>Ring A/B-monooxymonocyclic sterane? (C₂₃)</td>
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<td>378(28), 253(100), 196(35)</td>
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<td>378</td>
<td>—</td>
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<td>Ring A/B-monooxymonocyclic sterane? (C₂₃)</td>
<td>366</td>
<td>394(68), 255(50), 145(48), 81(100)</td>
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<td>Ring A/B-monooxymonocyclic sterane? (C₂₃)</td>
<td>406</td>
<td>406(48), 98(60), 97(100)</td>
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<td>48</td>
<td>Ring A/B-monooxymonocyclic sterane? (C₂₃)</td>
<td>392</td>
<td>252(50), 349(100), 235(48), 211(43)</td>
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<td>Alkylbenzene</td>
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<td>396(13), 381(5), 231(100)</td>
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<td>Isomer of 29 (3,4,7,12a-tetramethyl-1,2,3,4,4a,11,12,12a-octahydrocrysene)</td>
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<td>378(100), 253(50), 211(70), 159(45)</td>
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<td>Toluene</td>
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<td>Unknown</td>
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</tr>
<tr>
<td>57</td>
<td>Unknown</td>
<td>300</td>
<td>300(100), 150(20)</td>
</tr>
</tbody>
</table>
APPENDIX
Chemical Structures
(Numbers and letters refer to Tables 2 and 3.)

squalane (k ?)

diasterenes
R=H, CH₃, C₂H₅

ster-4-ene
R=H (g'), CH₃, C₂H₅ (g')

ster-5-enes
R=H (a), CH₃, C₂H₅ (a')

5α (H)-steranes
R=H (v), CH₃, C₂H₅ (v')

24-methylcholesta-4, 22-diene (w)

24-methylcholesta-5, 22-diene (x)

24-methylcholesta-5, 22-diene (x)

17α (H)-hopanes
R=H (y), C₂H₅ (y'), CH (CH₃)₂ (y')

17β (H)-hopanes
R=H (z), C₂H₅ (z'), CH (CH₃)₂ (z'), CH (CH₃) C₂H₅ (z'), CH (CH₃) C₃H₇ (z')

30-normotane (t')

hop-13 (18)-enes
R=H (h'), CH₃ (h')

hop-17 (21)-enes
R=H (g'), CH₃ (g')

fern-8-ene (k')

fern-9 (11)-ene (g')

fern-7-ene (p')

ionene (3)

2-methylbiphenyl (6)

phenanthrene (8)

norhydroabietane (16 ?)

pyrene (17)

27 ?

28

29 ?

35

43

C-ring monoarom. steroids
R=H, CH₃, C₂H₅

A-ring monoarom. steroids
R=H, CH₃, C₂H₅

anthrasteroids
R=H, CH₃, C₂H₅

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Plate 1. Petrography of organic matter, Leg 75 (Figs. 1, 3–5, and 8–11, transmitted light, ×160; Figs. 2, 6, 7, 12, normal reflected light, ×400).

1. Amorphous liptinite, dinoflagellate, humic matter; Sample 530A-97-3, 105–110 cm.
2. Bituminite II and frambooidal pyrite; Sample 530A-97-3, 105–110 cm.
3. Amorphous humic matter and pollen; Sample 530A-93-5, 135–145 cm.
6. Bituminite III with frambooidal pyrite; Sample 530A-103-3, 110–120 cm; separated kerogen.
7. Bituminite II and III in mineral matrix (may be degraded dinoflagellate? and spores) with frambooidal pyrite; whole rock block; Sample 530A-103-3, 110–120 cm.
9. Mixed organic facies. Amorphous humic and liptinite matter with spore; Sample 530A-105-1, 90–100 cm.
10. Amorphous humic and liptinite matter; Sample 532-20-2, 120–130 cm.
11. Mixed organic facies with zooclasts(?) and amorphous humic matter; Sample 532-7-3, 90–100 cm.
12. Bituminite II with frambooidal pyrite; Sample 532-37-2, 110–120 cm.