Article
Geochemical Characteristics and Significance of Organic Matter in Hydrate-Bearing Sediments from Shenhu Area, South China Sea

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Abstract: Rock-Eval pyrolysis and the biomarker composition of organic matter were systematically studied in hydrate-bearing sediments from the Shenhu area, South China Sea. The n-alkane distribution patterns revealed that the organic matter in the sediments appeared to originate from mixed sources of marine autochthonous input, terrestrial higher plants, and ancient reworked organic matter. The low total organic carbon contents (average < 0.5%) and the low hydrogen index (HI, <80 mg HC/g TOC) suggested the poor hydrocarbon-generation potential of the deposited organic matter at a surrounding temperature of <20 °C in unconsolidated sediments. The abnormally high production index and the fossil-originated unresolved complex mixture (UCM) accompanied by sterane and hopane of high maturity indicated the contribution of deep hydrocarbon reservoirs. Preliminary oil-to-source correlation for the extracts implied that the allochthonous hydrocarbons in the W01B and W02B sediments might have originated from the terrestrial source rocks of mature Enping and Wenchang formations, while those of W03B seem to be derived from more reduced and immature marine source rocks such as the Zhuhai formation. The results of the organic extracts supported the previous identification of source rocks based on the isotopic composition of C2+, hydrate-bound gases. The biomarker of methanogens, squalane, was recognized in the sediments of this study, possibly suggesting the generation of secondary microbial gases which are coupled with the biodegradation of the deep allochthonous hydrocarbons.

Keywords: organic matter; biomarkers; Rock-Eval; gas hydrate; biodegradation; South China Sea

1. Introduction

Natural gas hydrates are crystalline compounds composed of water and gases formed under high pressure and low temperature [1], which have been a frontier issue in both the industrial and academic research fields [2,3]. Marine geological surveys and drilling expeditions have shown that the Shenhu area, South China Sea (SCS) has abundant gas hydrate resources [4–7], and it has been one of the most promising exploration areas for gas hydrates [8–13].

It is conducive to understanding the formation mechanism, evolution history, and distribution of gas hydrate to acknowledge the origins and sources of hydrate-bound gas and its corresponding source rocks [3,14,15]. At present, most studies rely on the geochemical characteristics of gas hydrate, which provide limited available indicators
(mainly carbon and hydrogen isotopes of methane ($\delta^{13}$C-CH$_4$ and $\delta^D$-CH$_4$) and the dry coefficient ($C_1/(C_2 + C_3)$; $C_1$, $C_2$, and $C_3$ refer to methane, ethane, and propane, respectively) to study the gas origins [16–18]—the ambiguous boundaries in the genetic diagrams and the complex secondary processes present difficulties for a definite interpretation. Fortunately, biomarkers extracted from the hydrate-bearing sediment have more proxies and thereby provide more information regarding the original deposition and potential oil-source correlation of organic matter, and more clues as to the source rocks of the hydrate-bound gas.

The geochemical characteristics of gases bound in hydrates and core sediments at the three sites used in this study have been previously reported [19–22]. Hydrate samples from W03B were suggested to be dominated by microbial gases, while those from W01B and W02B exhibited a high content of thermogenic hydrocarbons [21,22]. Therefore, it is an intriguing phenomenon that these three drilling sites have different gas origins and sources, despite the fact that they are adjacent and located on either side of a canyon in the Shenhu area (Figure 1b). Moreover, for the Shenhu area, it has been reported that the deep conventional hydrocarbons have undergone an upward migration and subsequent microbial biodegradation to participate in the formation of shallower gas hydrate, based on the gas geochemistry and microbial community [21–23], but this inference still lacks further robust evidence. Remarkably, biomarker patterns are important evidence to shed new light on the contribution of petroleum hydrocarbons from deep oil/gas reservoirs and provide clues for potential oil/source correlation. Apart from this, lipid biomarkers in gas hydrate systems may provide important information for microbial communities associated with methane metabolic activities.

Figure 1. (a) Location of the Shenhu area in the Baiyun Sag of the Pearl River Mouth Basin (PRMB), South China Sea (SCS), modified from [24]; (b) location of the hydrate coring sites of this study—W01B, W02B, and W03B. C1–C17 refer to 17 canyons in the Shenhu area, modified from [10,21,25].
This study investigates the organic biomarker and pyrolysis geochemistry of sediment samples from three sites (W01B, W02B, and W03B) in the Shenhu area by Gas Chromatography–Mass Spectrometry (GC–MS) and Rock-Eval analyses to determine: (1) the sources of the organic matter in the sediments; (2) the microbial gas generation potential of the in situ sediment; (3) the existence/absence of a deep hydrocarbon contribution and potential oil-source correlation; and (4) the significances of organic matter for the gas hydrate system research in the Shenhu area. The novel results of this study might provide profound insights into gas hydrate formation in the Shenhu area.

2. Geological Setting

Located at the intersection of three tectonic plates, the SCS is the largest passive marginal sea in the Western Pacific [26,27]. The complex history of tectonic evolution and the promising prospects of oil and gas have made the SCS a natural laboratory for marine geology research [28,29]. The Shenhu drilling area is located in the Baiyun Sag, a deep-water depression in the Pearl River Mouth Basin (PRMB) of the SCS [30,31] (Figure 1), which has favorable conditions for gas hydrate formation and preservation [32–35]. High-quality source rocks have been found to supply gas for the shallow gas hydrate deposits [11,12,36–38]. The Eocene Wenchang formation and the Oligocene Enping formation are considered to be mature/overmature source rocks for the generation of thermogenic gases (Figure 2) [11,37,38]. The Zhujiang formation and the Hanjiang formation cannot provide thermogenic gas due to the limitation of their maturity, but they can still provide microbial gas as source rocks (Figure 2) [12,36]. Conventional oil/gas reservoirs have been explored in the PRMB, further confirming the great resource potential of this area [11,39–41]. Furthermore, polygonal faults induced by the overpressure environment of rapid deposition and large-scale mud diapirs formed during the neotectonic movement, all providing migration channels for gases accumulating at the gas hydrate stability zone (GHSZ) [32]. In addition, regional exploration shows that the Baiyun Sag has favorable geological conditions (seafloor pressures greater than 10 MPa, seafloor temperatures below 4 °C, and a geothermal gradient of 45–67.7 °C/km) for gas hydrate preservation [42,43]. Furthermore, it is noteworthy that the Shenhu area is the first place in the world where high concentrations of disseminated natural gas hydrate have been discovered in fine-grain sediment [9,25,44–46]. The gas hydrate is finely distributed in the foram-rich clayed silts, with a concentration of 20–60% of the pore volume [9,25,44].
Figure 2. Structural evolution characteristics and stratigraphic column of the PRMB, modified from [31,47].

3. Results

3.1. Rock-Eval Pyrolysis

The total organic carbon (TOC) contents of sediments from W01B and W02B decreased exponentially with depth from 1.34% to 0.15% (Figure 3a,e), while that of W03B slightly increased with depth from 0.19% to 0.32% (Figure 3i). The S₁ and S₂ of sediments from the three sites all showed extremely low values, and the depth variation profiles were similar to those of the TOC (Figure 3b,f,j). The HI values of sediments from the three sites ranged from 7 to 77, 16 to 55, and 16 to 45 mg HC/g TOC, respectively (Figure 3c,g,k). The OI indexes of sediments from the three sites were in the range of 40–125, 100–139, and 74–93 mg HC/g TOC, respectively (Figure 3c,g,k). The production indexes (PIs) of sediments from the three sites were in the range of 0.11–0.86, 0.15–0.58, and 0.11–0.5, respectively (Figure 3d,h,l). The T_max values of the sediments were in the range of 331–404 °C, 366–399 °C, and 374–405 °C, respectively (Figure 3d,h,l). A detailed data composition is shown in Table S1.

3.2. Biomarkers

3.2.1. n-Alkanes

The samples obtained from the three sites mainly contained n-alkanes from n-C₁₂ to n-C₃₅, and most of the samples displayed a bimodal distribution pattern (Figure 4) with a maximum at n-C₁₆ or n-C₁₈ of the pre-peak and n-C₂₉ or n-C₃₁ of the post-peak (Figure 4a). Most samples showed a bimodal distribution pattern with the pre-peak dominating (e.g., Figure 4b: 3-18X, 1-17X), except for the shallow samples, which showed the post-peak dominating (e.g., Figure 4b: 1-2H). The pre-peak showed an even-carbon dominance, while the post-peak showed an odd-carbon dominance (Figure 4a).
Figure 3. Depth profile of total organic carbon (TOC) (including pyrolysable organic carbon (PC) and residual organic carbon (RC)), free hydrocarbons ($S_1$), oil potential ($S_2$), hydrogen index (HI), oxygen index (OI), production index (PI), and $T_{\text{max}}$ of W01B (a–d), W02B (e–h), and W03B (i–l).
Area percentage

Abundance

Figure 3. Depth profile of total organic carbon (TOC) (including pyrolysable organic carbon (PC) and residual organic carbon (RC)), free hydrocarbons (S 1), oil potential (S 2), hydrogen index (HI), and oxygen index (OI), production index (PI), and Tmax of W01B (41.7 mbsf); W02B (99.9 mbsf); W03B (176.36 mbsf).

Figure 4. (a) Total ion chromatogram (TIC) of extracts from the studied sample W02B-4H (99.9 mbsf); (b) representative n-alkane composition from W01B-2H (54.9 mbsf), W01B-17X (166.88 mbsf), and W03B-18X (176.36 mbsf).

3.2.2. Isoprenoid

The two major acyclic isoprenoid hydrocarbons in all the samples were 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 [48,49]. The sources of pristane and phytane are multiple and complex, but, as a general rule, Pr/Ph values > 3 are typical of organic matter with predominantly terrestrial sources deposited in or transported through an anoxic environment; values of less than 1 are typically found in sediments from anoxic marine or hypersaline depositional environments [48,50,51]. The Pr/Ph of samples from the three sites ranged from 0.6 to 1.11, 0.61 to 0.97, and 0.18 to 0.8, respectively; all were less than 1 except for W01B-10x (1.11) and W01B-17X (1.04), indicating a phytane-dominant reduction depositional environment. Except for Pr and Ph, another kind of isoprenoid, squalene, was detected in samples from all three sites, of which the content ranged from 1.38 to 10.06, 2.49 to 7.84, and 3.71 to 13.19, respectively. The total squalane content of W03B was higher than that of W01B and W02B.

3.2.3. Steranes and Terpanes

A series of terpanes were detected in the extracts, including tricyclic terpanes (TT), tetracyclic terpanes (TeT), and hopanes series (Figure 5a). Figure 5a shows the distribution patterns of the hopane series. They are characterized by the predominance of C30 hopane (C30H) and C29H. The relatively high abundance of biological configurational isomers 17β(H), 21β(H)-hopane (ββ-C30H) and 17β(H), 21β(H)-homohopane (ββ-C31H) indicate their low maturation levels (Figure 5a). The 18α(H)-22,29,30trisnorhopane (Ts) had a relatively high abundance in all samples, with Ts/Tm ratios ranging from 0.25 to 0.81, 0.19 to 0.72, and 0.11 to 0.55, respectively. C31 22S/(S + R) showed less variation than Ts/Tm,
with values of 0.23–0.53, 0.21–0.4, and 0.1–0.21, respectively, whereas gammacerane (Ga), oleanane (OL), and C_{33}–C_{35} homohopanes were absent. C_{27} diasteranes and C_{27}, C_{28}, and C_{29} regular steranes were detected in the extracts (Figure 5b). The C_{27}, C_{28}, and C_{29}-5α(H),14α(H),17α(H)-20R-cholestanes exhibited an approximately V-shaped distribution pattern (Figure 5b). The sterane maturation parameter ααααC_{29}20S/20(S + R) index of the three sites ranged from 0.16 to 0.43, 0.15 to 0.36, and 0.03 to 0.24, respectively, and the C_{29}ββ/ββ + αα index of the three sites ranged from 0.24 to 0.42, 0.23 to 0.43, and 0.07 to 0.49, respectively.

Figure 5. (a) Distributions of terpane and hopanes on the mass chromatograms (m/z 191). Peak assignments: TT—tricyclic terpane; H—hopane; M—moretane; Ts—18α(H)22,29,30-trisnorhopane; and Tm—17α(H)-22,29,30-trisnorhopane. R, S: the configuration of the compounds. (b) Distributions of the diasteranes and regular steranes on the mass chromatograms (m/z 217).

4. Discussion

4.1. Sources and Preservation of Organic Matter

4.1.1. n-Alkanes

It is thought that the light hydrocarbons (C_{21}–) are mainly derived from the lipids of marine algae, while the long-chain hydrocarbons (C_{21}+) mainly originate from the waxes of terrestrial higher vascular plants and the lipid fragments derived from conifers [52–54]; thus, the bimodal distribution pattern of most of the samples in this study indicates a terrestrial–marine mixed source (Figure 4a,b), except for the layers such as W01B-10X and W01-17X, which tended to be unimodal pre-peak (Figure 4b), and layers such as W01B-2H and W02B-1H, which had an obvious unimodal post-peak (Figure 4b). The TAR, C_{21}/C_{21}+, and OEP2 showed that W01B and W02B had more terrestrial contributions in the shallow part (Table S1), while W03B had the opposite tendency, which may be related to the geographical location of the three sites. W01B, W02B, and W03B were located on either side of the migration canyon in the Shenhua area, respectively, and formed a unique sedimentary pattern under the complex interaction of a gravity current, a bottom current, and internal waves, resulting in differences in deposition between the sites [55–60].

In addition to the mixed-source characteristics, it was remarkable that C_{12}–C_{22} n-alkanes with even-to-odd predominance were observed in all samples (Figure 4). They were also observed in recent/ancient sediments as well as in the marine sediments [61–65]. The possible reasons for the even–odd dominance of short-chain n-alkanes (C_{12}–C_{22}) in the sediments are as follows: (I) normal fatty acids with odd-carbon predominance were reduced in a strongly reducing environment during deposition [66]; (II) anthropogenic pollution from fossil fuels or hydrocarbon leakage from the underlying high-maturity formations [67]; (III) biomass combustion can also form the dominant distribution of normal short-chain carbon pairs (thermal degradation of long-chain n-alkanes to short-chain n-alkanes) [68]; and (IV) they are derived from specific species of bacteria or fungi [62,
All the layers of the three sites displayed obvious even-carbon predominance, which is presumed to be independent of hydrocarbon leakage but usually indicates the direct input of microorganisms in sediments.

Compared to short-chain \( n \)-alkanes, long-chain \( C_{23} - C_{33} \) \( n \)-alkanes with a strong odd–even carbon preference (particularly \( C_{27}, C_{29}, \) and \( C_{31} \)) are indicative of terrestrially derived organic matter [71]. On the other hand, \( n \)-alkane distributions without odd/even predominance in sediments lacking petrogenic markers (hopane and sterane mixtures specific to mature organic matter) have been attributed to microbial odd-C-numbered inputs or to the microbial reworking of the \( n \)-alkane mixtures [70,72–74], suggesting that they can be used as tracers for degraded or microbially altered organic matter [75,76]. Both types of \( n \)-alkane distribution were quantified separately in the sediment from the three sites using the following equations [77,78]:

\[
A_{\text{odd}} = \sum_{i=1}^{6} C_{(21+2i)} - \sum_{i=1}^{5} C_{(22+2i)} - \frac{C_{22} + C_{32}}{2}
\]

(1)

\[
A_{\text{rew}} = C_{t} - A_{\text{odd}}
\]

(2)

where \( C_{i} \) is the measured concentration of each \( n \)-alkane and \( C_{t} \) is the sum of all \( n \)-alkane homologues between \( C_{21} \) and \( C_{33} \). \( A_{\text{rew}} \) refers to \( n \)-alkanes without an even/odd carbon number preference and \( A_{\text{odd}} \) refers to odd carbon number dominant \( n \)-alkanes.

From the calculation results, W01B and W02B were shown to have more contributions from terrestrial plants in the shallow part and more contributions from reworked organic matter in the deep part (Figure 6a,b). However, the proportion of W03B was relatively constant with depth and had the highest contribution from reworked organic matter among the three sites (Figure 6c). From the perspective of \( n \)-alkanes, organic matter from the Shenhu area appeared to be from mixed marine, continental, and ancient reworked organic matter sources, with the latter possibly contributed by turbidity flows/bottom currents from the upper continental shelf or nearby continental slope.

**Figure 6.** Relative proportion of \( n \)-alkanes without an even/odd carbon number preference (\( A_{\text{rew}} \)) and odd carbon number dominant \( n \)-alkanes (\( A_{\text{odd}} \)) of \( n \)-alkanes in sediment samples from (a) W01B, (b) W02B, and (c) W03B. Blank area and shaded area represent \( A_{\text{odd}} \) and \( A_{\text{rew}} \), respectively.

### 4.1.2. Steranes

Information regarding source type may sometimes be obtained from the relative abundance of \( C_{27} - C_{29} \) sterols or their fossil counterparts, the steranes [79,80]. 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 [50, 81, 82].

C_{27}/C_{29} indicates that the marine contribution of organic matter in W03B was higher than that of W01B and W02B. However, in some horizons, such as W01B, there was an increase in the pre-peak of the n-alkane distribution (low carbon number), but there was no corresponding reflection in the C_{27}/C_{29} indicators. In this case, we suggest that the n-alkane distributions better reflect the terrestrial influxes.

4.1.3. Pyrolysis Indexes

The Rock–Eval method developed by Espitalie et al. [83] is a widely used method for petroleum source-rock characterization and evaluation [84–86]. The HI, which correlates strongly with the elemental H/C ratio of the kerogen, is used to determine the hydrogen richness of a rock or sediment in order to identify the type of organic matter originally deposited in a given setting and to infer some aspects of the preservation of the remaining organic matter. High HI values are interpreted to reflect both algal and planktonic (high H/C) sources and the good preservation of the deposited organic matter [83], whereas low HI values are used to identify terrestrial organic matter sources and/or the extensive post-depositional alteration (oxidation) of algal remains [87]. Specifically, in immature marine sediments containing mixed phytoplankton, zooplankton, and bacterial debris (Type II organic matter), typical HI values range between 200 and 400 mg HC/g TOC [83], whereas in sediments containing terrestrial organic matter (Type III), the HI values are less than 200 mg HC/g TOC. Very low HI values are observed in sediments containing oxidized and reworked organic matter (Type IV) [85]. Consequently, the sediment records of HI provide information on both the sources and post-deposit processes of organic matter.

Cross plots of the HI, OI, and T_{\text{max}} of the sediment samples from the three sites are shown in Figure 6 [85]. These plots may be used similarly to a van Krevelen diagram to differentiate organic matter types. The samples in this study fell into the category of Type IV organic matter due to a lower HI (Figure 7a, b). A low HI index has been attributed to: I. A mineral matrix effect, if the sediment has variable amounts of carbonate and clay minerals, which will result in low but variable HI values [85, 88, 89]. However, the carbonate fraction was removed before the Rock-Eval analysis, and the clay mineral content was confirmed to be low and constant among the different samples in this study (unpublished data), so the mineral matrix effect was not considered to be the dominant factor for the low HI. II. Additional contribution from varying amounts of hydrogen-poor, reworked organic matter. Sediment reworking can increase the relative proportion of terrestrial organic matter by preferentially removing algal remains and lower the HI values by increasing the degradation degree of organic matter during post-depositional winnowing. The relatively lower HI/OI (<0.5) can also help to reconfirm this circumstance; a low HI and HI/OI in this study indicated a contribution from highly degraded reworked organic matter, since fresh or not highly degraded organic matter usually has a HI/OI greater than 2.

The low HI value is consistent with the previously calculated ancient reworked organic matter proportion in the samples. Specifically, the W03B site showed a higher proportion of reworked organic matter and also had the highest T_{\text{max}} and lowest HI values of the three sites (Figure 7b). The explanation for this phenomenon relies on the high proportion of reworked allochthonous organic matter, which manifested itself as a high T_{\text{max}} value and a low HI value, while the immature autochthonous component was responsible for the opposite signature.
I. Mineral matrix effect, if the sediment has variable amounts of other minerals, can influence the TOC content. This effect becomes more prominent in sediments with low TOC content (<0.4%), which may not reach the threshold of in situ methane generation and gas hydrate formation. Additionally, as mentioned before, very low HI values were observed in the sediments from this study, which indicated the presence of oxidized and reworked organic matter (Type IV) and corresponded to a poor hydrocarbon generation potential [85]. Moreover, according to the seafloor temperature (~4 °C) and geothermal gradient (54.6–62.6 °C/km) reported by Zhang et al. [21], the temperature of the studied layers (<240 m) was calculated to be less than 20 °C, which is below the optimum temperature zone for microbial activities. Furthermore, the pyrolysis parameters 

$$\text{T}_{\text{max}}, \text{S}_1, \text{S}_2, \text{HI}, \text{T}_{\text{max}}$$

and I of the sediment samples from the three sites had extremely low values (Figure 7), which were far lower than the threshold criteria for source rocks (Peters and Cassa, 1994). Therefore, such depleted TOC contents (most < 0.4%), in situ temperatures (<20 °C), and HI indexes (mostly less than 50 mg HC/g TOC) may jointly suggest a fairly poor hydrocarbon generation potential for the in situ sediments at these three sites that is insufficient to form highly saturated gas hydrate relying solely on the in situ sediment.

4.2. The Potential of Microbial Gas Generation in Hydrate-Bearing Sediments

In the diagenesis stage (also called the immature stage), the microbial gas generation potential of sediments is mainly controlled by the TOC content [90–93]. The formation of gas hydrate with low saturation (1–2%) from in situ sediments requires a minimum TOC content of 0.5% [93–95]. For the formation of high-saturation hydrate (>3%), it is not sufficient to rely solely on the biodegradation of particulate organic matter [96]. In addition, an optimal temperature (35–65 °C) is required during microbial methanogenesis activities for methanogens to survive and produce methane efficiently [90,93]. In a word, the high yield of microbial gas requires a high abundance of reactive/labile organic matter and a favorable temperature.

The TOC contents of the studied samples from the three sites were extremely depleted (average of 0.37% for W01B and W02B, 0.13% for W03B) and did not reach the threshold of in situ methane generation and gas hydrate formation. Additionally, as mentioned before, very low HI values were observed in the sediments from this study, which indicated the presence of oxidized and reworked organic matter (Type IV) and corresponded to a poor hydrocarbon generation potential [85]. Moreover, according to the seafloor temperature (~4 °C) and geothermal gradient (54.6–62.6 °C/km) reported by Zhang et al. [21], the temperature of the studied layers (<240 m) was calculated to be less than 20 °C, which is below the optimum temperature zone for microbial activities. Furthermore, the pyrolysis parameters 

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4.3. Implication of Upward Migration of Deep Hydrocarbons

4.3.1. Evidence of Petroleum Hydrocarbons

Primarily, the pyrolysis parameters 

$$\text{S}_1, \text{S}_2, \text{T}_{\text{max}}, \text{HI}$$

and I of the samples were significantly lower than the threshold for source rocks [97], indicating that thermogenic hydrocarbons could not be generated. However, most of the samples had a PI greater than 0.1, even as high as 0.8 (W01B-16, -17X; W02B-7H) (Figure 8d), indicating the intrusion and preservation of allochthonous hydrocarbons into the sediment [85].

In addition, an unresolved complex mixture (UCM) composed of a large number of isomers and homologues with complex branched chains and acyclic compounds that cannot be resolved by capillary columns was present in most samples, as shown in the chromatogram in Figure 4a [98,99]. The presence of a UCM in marine sediments is an
indicator of chronic/degraded petroleum contamination [100]. Although a bacteria-derived UCM cannot be ruled out [101], the GC/MS identification of 17α (H), 21β (H)-hopanes and steranes confirmed the fossil origin of the UCM (Figure 8a,b). Conventional oil and gas fields are ubiquitous in the northern SCS, and the hydrate gas has been considered to be cogenetic with the deep conventional gas reservoir; therefore the detection of the petroleum-based contaminants in this area was not unusual.

Moreover, the increase in the steranes and terpanes maturity index at the W01B and W02B sites also suggested the supply of deep mature hydrocarbons (Figure 8a,b [50,87,101–113]). The thermal effects of the stereochemical complexity of the basic skeleton of steranes and terpanes, i.e., the C29ββ/(ββ + αα), C2920S/(20S + 20R), Ts/Tm, and C31αβ22S/22(S + R), could help to determine the maturity and the cut-off points between the immature, low-mature, and mature stages, which are labeled in Figure 8a,b [51,81,102–114]. The sterane and terpane maturity parameters of the samples from sites W01B and W02B of this study indicated that the organic matter mainly originated from low-mature to mature source rocks.

![Figure 8](image-url)
4.3.2. Preliminary Oil-to-Source Correlation in the Extracts

Previous studies have suggested that the Baiyun Sag contains three sets of potential source rocks: Wenchang formation lacustrine source rocks, Enping formation transitional source rocks, and Zhuhai formation marine source rocks [11,21,41,116,120,121]. The first two sets of source rocks are identified as the principal source rocks, as they have a relatively high hydrocarbon generation potential and are mainly in the high-mature stage. The Zhuhai formation source rock is not considered as a primary source rock, because of its low thermal maturity [11,41,122,123]. The source rocks of the Wenchang formation Group formed in relatively anoxic to suboxic, lacustrine conditions, while the source rocks of the Enping formation were deposited in oxic, shallow lacustrine settings, with a relatively high input of terrigenous higher plants at the beginning and subsequent impact from a marine transgression; the source rocks of the Zhuhai formation were deposited in anoxic, relatively stable, neritic environments, with a greater contribution from aquatic algae relative to terrestrial higher plants [116,117].

To further investigate the origin of the allochthonous hydrocarbons within the extracts, a preliminary oil-to-source correlation was carried out based on the geochemical results obtained in this study and the geochemical parameters of the representative source rocks in the Baiyun sag reported in the literature. A series of biomarker compounds from the mature source rocks were also detected in the extracts, except for the immature hydrocarbons, such as the isoprenoids (Pr and Ph) on the mass chromatograms (Figure 4a), tricyclic terpanes and hopanes (Figure 5a), and diasteranes and regular steranes (Figure 5b). The distributions and characteristics of these biomarkers were useful for illustrating the origin of the allochthonous hydrocarbons within the extracts. In this study, Pr and Ph were detected in all samples, and the Pr/Ph ratios ranged from 0.18 to 1.11 (Figure 8c), suggesting that the organic matter sources of the allochthonous hydrocarbons were mainly deposited in an anoxic depositional environment [48,50,51]. The plot of Ph/nC\textsubscript{18} versus Pr/nC\textsubscript{17} (Figure 8d) shows that W01B and W02B had more terrestrial organic matter under more oxidized conditions, while W03B indicated more marine sources and a more reduced depositional environment. W01B and W02B exhibited geochemical characteristics similar to those of the terrestrial source rocks of the Enping and Wenchang formations, of which the latter contributed more to the supply. While W03B displayed geochemical characteristics similar to the marine source rocks of the Zhuhai formation.

Combined with the abnormally high maturity and the geochemical characteristics similar to those of deep mature source rocks, it is speculated that the allochthonous hydrocarbons within the extracts of W01B and W02B may include contributions from the Enping and Wenchang formations, of which the latter has a higher similarity, while W03B has a greater contribution from the immature marine source rocks of the Zhuhai formation.

4.4. Implication for Hydrate-Bound Gas Origin

4.4.1. Evidence of Thermogenic Gas Contribution

The combination of a poor microbial gas generation potential and a weak biodegradation degree both indicate that the in situ organic matter could not generate sufficient gases for the formation of hydrate with a saturation as high as 20–60% at the three sites [25], indicating that there must be other gas sources supplying the hydrate system in this area; this is also consistent with our above speculation as to deep hydrocarbon leakage.

From the perspective of gas geochemistry, it has been suggested that the hydrate-bound gases of sites W01B and W02B are mixed gases dominated by thermogenic gases [19–22], while the gas origin of W03B is mainly microbial gases produced by the biodegradation of organic matter in shallow strata (Figure 9a,b) [21], which further enhances the speculation that W01B and W02B received contributions from deeper thermogenic hydrocarbons. Liang et al. [22] obtained the isotope composition of the C2\textsubscript{4} hydrocarbons bound in hydrate in the Shenhu area to identify the source rocks of the gas hydrate and found that the biogenic hydrocarbons and thermogenic hydrocarbons were derived from marine organic matter and terrestrial organic matter, respectively. The source rocks of ther-
mogenic hydrate gas are interpreted to be from both the gas-prone coal strata of the Enping formation and the oil-prone medium-deep lacustrine strata of the Wenchang formation, the latter of which contributed more to the hydrocarbon supply of the gas hydrates [22]. These conclusions are consistent with the results of the oil-to-source correlation via organic biomarkers in this study. In addition, some studies show that the source rock of the Wenchang formation has been in the stage of pyrolytic dry-gas production since the early Pliocene [117,124] and is considered to be the main source rock for hydrate accumulation in this region, which is consistent with the conclusion drawn as to the extract sources in this study and the gas sources in the previous study.

4.4.2. Evidence of Secondary Microbial Gas

Squalene was detected in this study, which has been used as a biomarker for methanogens [48]. A study by Lin et al. [23] verified that most abundant archaeal genera are potential methanogenic archaea in the Shenhu area, and the hydrogenotrophic type was inferred to be predominant among the multiple types of methanogenesis. It is well-known that microbial biodegradation is coupled with the methanogenic process, which means that the presence of the methanogenic process also indicates the existence of biodegradation [100,125,126]. Considering the low hydrocarbon-generation potential and intact n-alkane preservation of the in situ sediment, we speculate that the methanogenesis activities are likely related to the biodegradation of deep allochthonous hydrocarbons, which is coupled with the generation of secondary microbial gases [127]. This inference corresponds to the speculation of Zhang et al. [21] that the geochemical characteristics of the hydrate-bound gas from these three sites are very similar to those of confirmed biodegraded gases [21,128,129], which is also consistent with the identified putative hydrocarbon degraders in the microbial community structure study by Lin et al. [23]. Moreover, the deviations in the temperature of methanogens from the geothermal gradient of the study area suggested the migration of organic matter, emphasizing the possibility of microbiobly mediated secondary microbial gas formation from deep thermogenic fluid [23]. Therefore, the gases bound in hydrate from the three sites may have complex sources, including the primary in situ microbial gases, deep thermogenic gases, and secondary microbial gases generated by the biodegradation of the leakage hydrocarbons.

Figure 9. Genetic diagram of (a) δD-CH4 versus δ13C-CH4 and (b) C1/(C2 + C3) versus δ13C-CH4 (the boundary lines are from [130]), showing the genetic type of the hydrate-related gas from sites W01B, W02B, and W03B. Data from [19-22]. A—abiotic; F—fermentation; CR—CO2 reduction; EMT—early-mature thermogenic; LMT—late-mature thermogenic; SM—secondary microbial.
5. Materials and Methods

The samples were retrieved by the GMGS4 expedition, which was carried out on the geotechnical drilling vessel Fugro Voyager in 2016 [25]. The water depth of the three sites is ~1285 m, 1274 m, and 1310 m, respectively, and the drilling depth was ~234 m below seafloor (mbsf), 240 mbsf, and 222 mbsf, respectively. Detailed information and lithological description of sediment samples can be found in Table S1.

5.1. Rock-Eval Analysis

Rock-Eval analysis was conducted by Rock-Eval IV instrument to obtain the amount of free hydrocarbons ($S_1$), the amount of hydrocarbons generated through thermal cracking of nonvolatile organic matter ($S_2$), the amount of CO$_2$ produced during pyrolysis of kerogen ($S_3$), and the temperature at which the maximum release of hydrocarbons from cracking kerogen occurred during pyrolysis (top of $S_2$, T$_{max}$). The hydrogen index (HI) was calculated from the formula HI = 1000 $\times$ $S_2$/C$_{org}$, and oxygen index (OI) was calculated using the formula OI = 1000 $\times$ $S_3$/C$_{org}$. The pyrolysis temperature of organic matter ranged from 300 °C to 600 °C, with a rising rate of 25 °C/min.

The Rock-Eval analyses were conducted at the National Research Center for Geoanalysis, China Geological Survey (CGS).

5.2. Biomarker Analysis

Around 100–200g dried and crushed samples (<100 mesh) were Soxhlet extracted for 72 h by the solvent mixture of dichloromethane (DCM) and methanol (93:7, v:v). After removing the solvents and asphaltenes by n-hexane, the resultant soluble fraction was separated into aliphatic hydrocarbons, aromatic hydrocarbons, and polar compounds by using the column chromatography method (alumina/silica gel column). Aliphatic hydrocarbons, aromatic hydrocarbons, and nonhydrocarbons were extracted by washing with n-hexane, dichloromethane, and methanol, respectively.

Aliphatic hydrocarbons were analyzed using a Gas Chromatography–Quadrupole Mass Spectrometer (GC–MS) 6890N/5973N (Agilent Technologies, Palo Alto, CA, USA), which was fitted with a DB-5 MS fused silica capillary column (J&W Scientific, Agilent, USA; 30 m $\times$ 0.25 mm $\times$ 0.25 µm). The GC oven temperature was initially set at 80 °C (hold for 5 min), then programmed to 290 °C (hold for 40 min) at a rate of 4 °C/min. The compounds were identified by comparing their mass spectra with those in the NIST02 library and published data.

The biomarker experiments were conducted in the sample pretreatment laboratory of the Key Laboratory of Petroleum Resource Research, Northwest Institute of Eco-Environment & Resources, Chinese Academy of Sciences.

6. Conclusions

The organic geochemical characteristics of hydrate-bearing sediment provide clues for studying the origins and formation processes of gas hydrate. In this study, the Rock-Eval pyrolysis and biomarkers (n-alkane, isoprenoid, sterane, and hopane) of organic matter were systematically studied in hydrate-bearing sediments from the Shenhu area (GMGS4-W01B, -W02B, and -W03B), South China Sea. According to the experimental results, the following conclusions can be drawn.

1. The n-alkane distribution patterns reveal that the organic matter in sediments from the three sites appears to originate from the mixed sources of marine, terrestrial higher plant, and ancient reworked organic matter; the presence of the latter is verified by the pyrolysis parameters, especially the low hydrogen index (HI) and high T$_{max}$. The relative proportion of terrestrial higher plants to reworked organic matter was calculated from equations, and it was found that the depth variation trend of the ratio was consistent with that of the HI and T$_{max}$, and the formation with a high proportion of reworked organic matter manifested as a low HI and high T$_{max}$.
(2) The low total organic carbon (TOC) contents (average < 0.5%) with a HI < 80 mg HC/g TOC suggest the poor hydrocarbon-generation potential of the deposited organic matter at a surrounding temperature of <20 °C in unconsolidated sediments, which also indicates that the highly saturated hydrate could not have been formed solely by the in situ sedimentary organic matter.

(3) The abnormally high production index (PI), fossil-originated UCM, and sterane and hopane of high maturity indicate the contribution of deep hydrocarbon reservoirs, which is consistent with the thermogenic origin contribution of hydrate-bound gases.

(4) The preliminary oil-to-source correlation for the extracts implied that the allochthonous hydrocarbons in the W01B and W02B sediments might have originated from the low-mature to mature terrestrial source rocks of the Enping and Wenchang formations, while those of W03B seem to be derived from more reduced and immature marine source rocks, such as those of the Zhuhai formation. The results of the organic extracts indicated that some organic matter might have migrated from deep hydrocarbon reservoirs, similar to the hydrate-bound gas, and supported the previous identification of source rocks based on the isotopic composition of C2+.

(5) That the biomarker of methanogens, squalane, was recognized in the sediments of this study might suggest the generation of secondary microbial gases which are coupled with the biodegradation of the deep allochthonous hydrocarbons, indicating the complex component of the hydrate-bound gases.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/molecules27082533/s1, Table S1: Site information and lithological description of the samples in this study; Table S2: Biomarker composition and Rock-Eval pyrolysis parameters of sediments in this study.

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