Molecular Characterization of NSO Compounds and Paleoenvironment Implication for Saline Lacustrine Oil Sands by Positive-Ion Mass Spectrometry Coupled with Fourier-Transform Ion Cyclotron Resonance Mass Spectrometry
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ABSTRACT: NSO compounds mainly exist in geological bodies in the form of nonhydrocarbons and asphaltenes with abundant geological and geochemical information. Combined with the gas chromatography mass spectrometry (GC–MS) technology, positive-ion electrospray ionization Fourier-transform ion cyclotron resonance MS (FT-ICR MS) was used to understand the composition and distribution characteristics of NSO compounds in the oil sands of the Dongpu Depression and to explore their paleoenvironmental significance. The results show that n-alkanes are characterized by an even carbon number and phytane dominance, suggesting a saline lacustrine environment. Certain abundance of nC37 and β-carotane, high gammacerane content, and low diasterane content are detected in the analyzed samples, reflecting the saline-reducing paleoenvironment for the organic matter. Nine types of heteroatom compounds are detected: N1, N1O1, N1S1, O1, O1S1, O2, O2S1, S1, and S2. The main compounds are S1 and N1 compounds, followed by O1S1 compounds. The double-bond equivalent (DBE) value of S1 compounds is mainly distributed between 3 and 12, and the carbon number is mainly distributed between 18 and 35. The DBE value of N1 compounds is mainly distributed between 4 and 14, and the carbon number is mainly distributed in the range 15–35. Among the S1 compounds, DBE3 compounds (thiophenes) have relatively more sulfur-containing carotenoids (C40). The abundance of C40 S1 and the ratio of pyridine and its homologue DBE4−8/DBE9−12 N1 compounds show a good contrast with the paleoenvironment indicators of gammacerane/C30 hopane and diasterane/regular sterane. They can be used as a reference for the paleoenvironment index. Maturity is another factor affecting the distribution of NSO heteroatoms in the oil sands. NSO compounds are enriched in the DBE area with higher condensation, and the main peak carbon shifts forward. As the maturity increases, the relative abundance of N1 compounds increases, the aromatization intensifies, and carbon is broken into short chains. The research results shed light on the potential application of NSO compounds in petroleum exploration based on FT-ICR MS.

1. INTRODUCTION
Oil sands are sedimentary rocks that contain hydrocarbons such as heavy oil, solid asphalt, and light oil.1 As an important alternative energy resource for oil and natural gas, oil sands are one of the most important strategic supplements to petroleum resources.2 Oil sands contain a large amount of nonhydrocarbons and asphaltenes where abundant NSO heteroatom compounds can be detected.3−5 NSO heteroatom compounds carry rich geological and geochemical information, such as geochemical origins,3−6 thermal maturity,7−12 and biodegradation.13,14 Understanding the molecular characteristics of NSO compounds in oil sands is of great significance for further studying the characteristics of oil sands and evaluating the resource potential. However, NSO heteroatom compounds are highly polar compounds with usually low amounts and, therefore, difficult to separate and identify. Conventional technical methods often fail to obtain comprehensive information on heteroatom compounds. At present, the geochemical studies of the NSO compounds are far fewer than those of conventional hydrocarbon compounds. Fourier-transform ion cyclotron resonance mass spectrometry (FT-
ICR MS is a new technique introduced in the field of petroleum geochemistry in recent years. It has ultra-high resolution and precision and can realize the identification of heteroatomic compounds at a molecular level. Because of the unique advantages for analyzing heteroatom compounds, using the electrospray ionization (ESI) FT-ICR high-resolution MS technology to study NSO compounds in oil sands is one of the research hot spots in the field of petroleum exploration. Previous studies showed that the distribution characteristics of NSO compounds of oil sands in different areas are quite different, and the effects of maturity and biodegradation affect distributions. However, some related factor indexes are mainly based on the analysis technology of negative-ion ESI, and the correlation between positive-ion ESI-related indicators and conventional biomarker parameters has rarely been reported yet.

The Dongpu Depression in the Bohai Bay Basin is one of the most typical saline lacustrine hydrocarbon-generating depressions in China. Despite its high degree of exploration, the composition and distribution of heteroatomic compounds in the oil sands are unclear, which hinders further exploration in this area. Therefore, in this paper, the FT-ICR MS technique is adopted combined with gas chromatography MS (GC–MS) to reveal the distribution characteristics of heteroatomic compounds in saline lacustrine oil sands and explore their geochemical significance, providing geochemical information for future oil and gas exploration in this area.

2. GEOLOGICAL SETTING

Located in the southeast corner of the Linqing Depression, the Dongpu Depression is an important part of the petroliferous Bohai Bay Basin (Figure 1a,b). It is bounded by the Lanliao Fault in the east and the Changyuan Fault in the west (Figure 1c). The southern part of the depression is separated by the Lankao Uplift, and it is connected to the Kaifeng Depression. The north of the depression is separated by the Maling Fault and is connected to the Xinxiang Depression in the Linqing Depression (Figure 1c). The overall shape of the depression is narrow in the north and wide in the south, with a north–northeast trend (Figure 1c). The Dongpu Depression is based on the Mesozoic and Paleozoic strata, with huge salt-bearing clastic rocks deposited in the Cenozoic (Figure 2). The Cenozoic strata developed the Paleogene Shahejie Formation and the Dongying Formation, and the Neogene developed the Guantao Formation and the Minghuazhen Formation (Figure 2). The Quaternary was deposited in the Pingyuan Formation with a total thickness of about 8000 m. Among them, the Shahejie Formation can be divided into Es4, Es3, Es2, and Es1; Es3 is subdivided into lower, middle, and upper members of Es3, with a stratum thickness of about 1500 m (Figure 2). The details of the stratigraphic characteristics are shown in Figure 2. The Paleogene is composed of the Shahejie Formation and the Dongying Formation, and the Neogene developed the Guantao Formation and the Minghuazhen Formation (Figure 2). The Quaternary was deposited in the Pingyuan Formation with a total thickness of about 8000 m. Among them, the Shahejie Formation can be divided into Es4, Es3, Es2, and Es1; Es3 is subdivided into lower, middle, and upper members of Es3, with a stratum thickness of about 1500 m (Figure 2). The details of the stratigraphic characteristics are shown in Figure 2. The Paleogene is composed of the Shahejie Formation and the Dongying Formation, and the Neogene developed the Guantao Formation and the Minghuazhen Formation (Figure 2). The Quaternary was deposited in the Pingyuan Formation with a total thickness of about 8000 m. Among them, the Shahejie Formation can be divided into Es4, Es3, Es2, and Es1; Es3 is subdivided into lower, middle, and upper members of Es3, with a stratum thickness of about 1500 m (Figure 2). The details of the stratigraphic characteristics are shown in Figure 2.
The lithology of each section from north to south varies greatly.26,27 According to its lithology and fossil combination, the Shahejie Formation is divided into four sections. The first member is about 250–450 m. The gypsum-salt rocks are mainly distributed in the northern area.26,29

### 3. MATERIALS AND METHODS

Five oil sand samples of different depths and layers were acquired from two wells, W18-5 and W79-8, in the Puwei area of the northern Dongpu Depression (Figure 1c). Three of them are from well W18-5, and the other two samples are from well W79-8. The sample experiment was carried out in the State Key Laboratory of China University of Petroleum (Beijing).

The oil sand sample was manually crushed, and 150–200 g was taken for Soxhlet extraction. Asphaltene is precipitated by n-pentane in the oil sand extract and saturated hydrocarbons. Hydrocarbon and nonhydrocarbon components are leached with petroleum ether (60 mL), dichloromethane (about 40 mL), and ethanol (30 mL). The extracts were separated and subjected to GC–MS.

#### 2.1. Whole Oil Analysis

Whole oil analysis was carried out using an Agilent 5890 gas chromatograph equipped with a DB-5 column (30 m × 0.25 mm i.d. and film thickness: 0.25 μm) with H2 as the carrier gas. The oven temperature was programmed to range from 40 to 310 °C (held 19 min) at 6 °C/min.

#### 2.2. GC–MS Analysis

The extracted saturated hydrocarbons were subjected to GC–MS analysis in the State Key Laboratory of China University of Petroleum (Beijing). GC–MS analysis of the hydrocarbon fractions of the samples isolated by column chromatography was conducted using an Agilent 6890 gas chromatograph with helium as the carrier gas. The oven temperature for the GC–MS analysis was set to 60 °C for 1 min, then increased to 120 °C at a rate of 4 °C/min and to 300 °C at 2 °C/min, and finally held at 300 °C for 20 min.

#### 2.3. Positive-Ion ESI FT-ICR MS

Bitumen (200 mg) was diluted with 20 mL of dichloromethane (CH₂Cl₂). A total of 50 μL of methyl iodide and 2 mL of silver tetrafluoroborate in 1,2-dichloroethane (20 mg·mL⁻¹) were added to the bitumen solution. The mixture in a beaker was immersed in an ultrasonic bath for 5 min and allowed to react at room temperature for 48 h. The precipitate of silver iodide was removed by centrifugation and then rinsed with CH₃Cl₂. The methyl thiophenium salts and the unreacted oil were obtained by evaporating CH₃Cl₂ from the centrifuged solution. A total of 5 μL of hexane was added to the mixture, and most of the unreacted oil was dissolved in the mixture and separated from the methyl thiophenium salts. Thiophenium salts (10 mg) were diluted with 1 mL of CH₃Cl₂. A total of 5 μL of a thiophenium salt solution was diluted with 1 mL of a toluene/methanol/CH₃Cl₂ (3:3:4) solution.

After the positive-ion methyl derivatization, the 9.4T FT-ICR MS instrument produced by Bruker was used, and the ESI positive-ion mode was selected. The injection speed is 150 L/h, the nozzle voltage is 2.5 kV, and the capillary inlet and outlet voltages are 3.0 kV and 320 V, respectively. The ion source hexapole DC voltage is 2.4 V, the radio frequency is 300 °C/min, and the ion storage time is 0.01 s. The ideal mass of Qₖ is 250 Da. The operating conditions of the quadrupole collision cell are 5 MHz, 400 Vp-p, and the ion storage time is 1 s. The transition time by electrostatic focusing for ion conversion into ICR is 1.3 ms. The working conditions of ICR MS include an

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**Figure 2.** Stratigraphic column showing the nomenclature, thickness, and lithology of major rock units of the Dongpu Depression.
excitation attenuation of 11.75 dB, a mass range of 200−750 Da, a data volume of 4M, and the time-domain signal superimposition 64 times. For data processing and analysis, see the literature.9,14

4. RESULTS AND DISCUSSION

3.1. Hydrocarbon Characterization by GC−MS.

3.1.1. Whole Oil Analysis. The total-ion chromatogram of the selected sand extracts is shown in Figure 3. The n-alkanes for the analyzed samples show even-carbon-number dominance with the carbon number range of 16−40. The carbon preference index (CPI) and odd−even predominance (OEP) values range from 0.86 to 1.02 and 0.84 to 1.02, respectively (Table 1), suggesting some maturity difference. Isoprenoids are rich in bitumens with a strong phytane advantageous distribution (Figure 3), suggesting a saline lacustrine environment. The Pr/Ph ratio ranges from 0.13 to 0.3 with an average of 0.19 (Figure 3, Table 1), suggesting its strongly reducing environment for the organic matter. The ratios of Pr/nC17 and Ph/nC18 did not have large differences between the samples, indicating good preservation and low or no biodegradation. A large amount of β-carotene was also detected in the bitumens, and the n-alkanes were rich in C37 and nC38 (Figure 3). Studies have shown that the predominance of nC37 is mainly controlled by the nature of the depositional environment.30 The high salinity and strongly reducing depositional environment are responsible for the abundance of nC37 in the geological body. In addition, β-carotene (Figure 3), which is commonly found in geological samples, usually occurs in an anoxic and strongly reducing saline environment.31,32 Ding et al. showed that β-carotene is mainly derived from the

Table 1. Geochemical Parameters of the Analyzed Samples

| well | sample no. | W18-5 | W18-5 | W18-5 | W79-8 | W79-8 |
|------|------------|-------|-------|-------|-------|-------|
| depth (m) | 2657.55 | 2715.39 | 2904.01 | 3363.96 | 3456.16 |
| Strata | Es3M | Es3L | Es3L | Es3L | Es3L |
| CPI | 0.86 | 0.95 | 0.95 | 0.88 | 0.88 |
| OEP | 0.84 | 0.92 | 0.94 | 0.90 | 0.90 |
| Pr/Ph | 0.16 | 0.18 | 0.16 | 0.13 | 0.30 |
| Pr/nC17 | 1.03 | 0.75 | 0.92 | 1.30 | 0.54 |
| Ph/nC18 | 2.73 | 1.75 | 1.95 | 2.79 | 1.22 |
| C21−22/C28−29 | 0.79 | 0.83 | 0.87 | 0.71 | 1.07 |
| ∑nC21−22/∑nC22−24 | 0.35 | 0.29 | 0.24 | 0.14 | 0.38 |
| 20S | 0.34 | 0.36 | 0.33 | 0.52 | 0.43 |
| αββ | 0.30 | 0.30 | 0.28 | 0.40 | 0.48 |
| Tc/(Tc + Tm) | 0.21 | 0.22 | 0.20 | 0.17 | 0.34 |
| dia/reg | 0.13 | 0.16 | 0.16 | 0.17 | 0.31 |
| ste/hop | 1.85 | 1.47 | 1.70 | 1.39 | 1.67 |
| G/C30H | 0.76 | 0.67 | 0.63 | 0.97 | 2.52 |
| C35H/C40H | 1.67 | 1.56 | 1.49 | 1.63 | 0.83 |
| DBTs (%) | 20.13 | 8.92 | 6.86 | 8.37 | 4.66 |
| C40 DBE, S, N1 | 6.09 | 5.63 | 6.66 | 4.78 | 2.55 |
| DBE1−4/DBE>15, N1 | 1.82 | 1.24 | 1.37 | 1.03 | 0.51 |

Note: CPI = [(nC23 + nC25 + nC27) + (nC25 + nC27 + nC29)]/2 × (nC24 + nC26 + nC28)/[(nC26 + nC28 + nC29)]; OEP = 1/[4(nC25 + 6nC27 + nC29)/[(nC26 + nC28 + nC29)]; 20S = C2020S/20S + 20R; αββ = C29 hydrocarbon/(ααα + αββ); dia/reg = C29 diasterane/sterane; ste/hop = sterane/hopane; G/C30H = gammacerane/C30 hopane; and DBTs (%) = the content of dibenzothiophene.
The selected samples are rich in C27, C28, C29 steranes, and the isomerization is relatively low.32 The terpanes for the analyzed samples show typical characteristics of a saline lacustrine environment. Algae.32 The ratio of gammacerane/C30 hopane ranges from 0.63 to 2.52, with an average of 1.11 (Table 1). Gammacerane is reported to be derived from photosynthetic bacteria and is enriched in crude oil and source rocks from saline lacustrine facies.32,34 The high abundance of gammacerane is usually associated with brackish water stratification, which shows that the organic matter was originated from the saline and reductive sedimentary environment.

### 3.1.2. Saturate Biomarker Analysis

The biomarkers sterane and terpane are detected in the oil sand extract, and representative samples of m/z 217 and m/z 191, respectively, are shown in Figure 4. The selected samples are rich in C27, C28, C29 regular steranes, and the isomerization is relatively low (Figure 4 left). The isomerization ratios of C29 acaaterane 20S/(S + R) and C29 aββα/(aββα + aααα) range from 0.28 to 0.48 and 0.33 to 0.52, respectively (Table 1), suggesting that the oil sands are at less mature to mature stages. The sample of B2 with a deeper depth corresponds to the equilibrium value, proving more mature than other samples (Table 1). The content of diasteranes is relatively low, and the regular sterane is relatively abundant (Figure 4 left). The ratio of diasterane/regular sterane ranges from 0.13 to 0.31 (Table 1). Studies have found that the formation of diasterane is related to the redox properties of the deposition environment.32,35 In an anoxic and strongly reducing environment with a Pr/Ph of less than 0.5, less diasterane is formed as the carbon skeleton of acidic diasterane is inhibited.33 Therefore, the low abundance of diasterane and the low ratio of diasterane/regular sterane are also signals of a strongly reducing paleoenvironment for the organic matter. However, there is an outlier diasterane/regular sterane of B2, which shows a higher ratio of diasterane/sterane than others (Table 1). Such cases were also reported in the reducing carbonates which depend on the amount of clay relative to total organic carbon.32 High diasterane/sterane ratios can result from high thermal maturity as well. Here, in the study area, we deduce that it is the maturity that should be responsible for the higher diasterane/sterane ratio of B2. The sterane/hopane ratios range from 1.39 to 1.85 (Table 1), indicating major contributions from planktonic and/or benthic algae.32 The terpanes for the analyzed samples show typical characteristics of a saline lacustrine environment. Abundant hopanes were detected, with C30 hopane as the main peak (Figure 4 right). High content of gammacerane is notably recognized for the analyzed samples. The ratio of gammacerane/C30 hopane ranges from 0.63 to 2.52, with an average of 1.11 (Table 1). Gammacerane is reported to be derived from photosynthetic bacteria and is enriched in crude oil and source rocks from saline lacustrine facies.32,34 The high abundance of gammacerane is usually associated with brackish water stratification, which shows that the organic matter was originated from the saline and reductive sedimentary environment.

### 3.2. Characteristics of NSO Compounds Analyzed by FT-ICR MS

#### 3.2.1. General Distribution Characteristics of NSO Compounds

The positive-ion ESI FT-ICR MS detected nine types of NSO heteroatom compounds in the oil sands of the Dongpu Depression, including N1, N1O1, N1S1, O1, O1S1, O2, O2S1, S1, and S2 (Figure 5). S1, O1S1, and N1 are the main ones, and their relative abundance distributions range from 25.17 to 49.82%, 6.73 to 37.26%, and 13.99 to 44.64%, respectively (Figure 5, Table 2), followed by N1,N1O1,N1S1,O1,O1S1, with their relative abundances in the ranges 3.36–44.64 and 2.71–5.89%, respectively (Figure 5, Table 2). Other types of compounds have lower abundances, usually less than 2% (Figure 5, Table 2). The N1 groups detected by positive-ion ESI FT-ICR MS are basic nitrogen groups such as pyridine, acridine, or quinoline.32 It is discovered that with the increase in maturity, the relative content of N1 compounds increased significantly, from 13.99 to 44.64% (Figure 5, Table 2), indicating the maturity effect on nitrogen compounds.
Figure 5. Relative abundance of NSO compounds in oil sands of the Dongpu Depression by ESI (+) FT-ICR MS.

Previous studies have found that nitrogen compounds are sensitive to maturity.35,36 With the increasing thermal maturity, the contents of various nitrogen compounds in the rock extracts increase drastically, together with significant compositional variations related to alkyl substitution.37 Although the nitrogen-containing compounds detected by positive ions are basic groups, they are controlled by their maturity and show consistency with the nonbasic "pyrrole-like" nitrogen compounds. The types of NSO compounds detected in the oil sands of the Dongpu Depression are in line with other marine basins or saline facies bodies such as the Tarim Basin,38 the Alberta area of Canada,39 and the continental basins developed with saline facies gypsum salt rocks such as the Jianghan Basin40 and Jinxian Depression.41 They are all rich in S1 compounds. However, the relative abundance differences of O1S1 and S2 are different. The compositional differences of NSO compounds may be related to their original source or depositional environment.

3.2.2. S1 Class Species and Its Geochemical Significance. The S1 class species is the most abundant species in the sample with a wide range of double-bond equivalent (DBE) and carbon number values. The carbon number and DBE composition and abundance of S1 compounds in the oil sands of the Dongpu Depression are shown in Figure 6. The carbon number of S1 ranges from 11 to 50, mainly distributed between 20 and 40 (Figure 6a–e), and the DBE value ranges from 0 to 23, mainly distributed between 4 and 15 (Figure 6f–j). As the maturity increases, the S1 compounds tend to be enriched in the area with high DBE values. The DBE values in the samples of A1–A3 are mainly distributed in the range 4–12 with the main peaks of DBE being 4 and 6, and the carbon number is mainly distributed in the range 15–35 (Figure 6a–c,f–h), while the DBE values of B1–B2 samples with higher maturity are mainly distributed in the ranges 6–12 and 6–20 with DBE values of 6 and 9 and 12 as the main peaks, respectively, and the carbon number distribution range is much wider within 16–40 (Figure 6d,e,i–j).

It is reported that the DBE value of 0 in the S1 class species is for alkyl sulfides, while DBE values of 1 and 2 for the S1 class refer to sulfides with one and two cyclic rings, respectively. Thiophenes have a DBE value of 3.9 Compounds of DBE values 6, 9, and 12 for the S1 class are for benzothiophene, dibenzothiophene, and alkyl dibenzothiophene, respectively.9 The benzo homologue of tetrahydrobenzothiophenes (one aromatic and one naphthenic ring) or phenylthiophene was identified for the DBE value of 7 of the S1 class. DBE values of 9 and 10 of the S1 class were for the benzo homologues with DBE values of 6 and 7, respectively.9 As shown in Figure 6, samples with lower maturity for A1–A3 contain less sulfides but higher abundance of benzothiophene compounds (Figure 6a–c,f–h). Benzothiophene (DBE6 S1) and dibenzothiophene (DBE9 S1) are the most abundant compounds, while sulfides and thiophenes are less abundant in the more mature samples of B1 and B2 (Figure 6d,e,i,j).

The carbon number distributions of DBE values 1, 3, 6, and 9 series compounds are shown in Figure 7. The carbon number of sulfide compounds (DBE1) ranges from 20 to 33. The carbon number range of samples A1–A3 with lower maturity is much narrower within 15–30, with C20 and C25 as the main peaks, followed by C30, C32, and C33 compounds (Figure 7). However, the distribution of B2 with higher maturity is quite different from others. The carbon number is mainly distributed between 16 and 22, with C17 and C19 as the main peaks (Figure 7a). The carbon number of thiophene species (DBE3) is mainly distributed between 22 and 33, with C24 and C25 as the main peaks (Figure 7b). In addition, thiophenes (DBE3) contains abnormally high C40 and relatively abundant C17 (Figure 7b). The carbon numbers of benzothiophenes and dibenzothiophenes (DBE6 and DBE9) are mainly distributed in the range of 20–36, with a certain abundance of C40 (Figure 7c,d). The more mature sample B2 has a lower abundance in the diabenzothiophene species with the main peaks of C14 and C15 (Figure 7d).

C40 S1 compounds with various DBE values having high relative abundance are likely to be sulfurized carotenoids that have been found in crude oils.42 Lexane (C33) was found to be abundant in the oil, which is also present in the desulfurized sulfur fraction (Figure 7b). It is reported that lexane is an important intermediate in the anaerobic thermal degradation of β-carotene.43 Sulfurized carotenoid C40 S1 compounds with various DBE values are good indicators of the original depositional environment. β-Carotene is highly specific for lacustrine deposition. Under highly reducing conditions, the carbon skeleton of carotenoids exists in the form of β-carotene, associating primarily with anoxic, saline lacustrine or highly restricted marine settings where organisms thrive as the dominant biota.11,12 Studies have shown that both sulfur complexes and kerogen in high-sulfur crude oil can release β-

Table 2. Relative Abundance of NSO Compounds Detected by Positive-Ion FT-ICR MS

| sample no. | depth (m) | Strata | N1 (%) | N2O1 (%) | N1S1 (%) | O1 (%) | O1S1 (%) | O2 (%) | O2S1 (%) | S1 (%) | S2 (%) |
|-----------|----------|--------|--------|----------|----------|--------|----------|--------|----------|--------|--------|
| A1        | 2657.55  | Es3M   | 13.99  | 4.53     | 0.16     | 2.55   | 21.49    | 0.48   | 5.84     | 49.82  | 1.14   |
| A2        | 2715.39  | Es3L   | 17.04  | 3.36     | 0.61     | 4.34   | 37.26    | 0.34   | 4.50     | 31.32  | 1.22   |
| A3        | 2904.01  | Es3L   | 21.73  | 4.40     | 0.78     | 4.22   | 34.38    | 0.74   | 5.91     | 26.61  | 1.23   |
| B1        | 3363.96  | Es3L   | 23.69  | 4.02     | 0.64     | 3.44   | 17.23    | 0.66   | 2.71     | 46.08  | 1.54   |
| B2        | 3456.16  | Es3L   | 44.64  | 13.21    | 0.39     | 1.46   | 6.73     | 1.51   | 5.89     | 25.17  | 0.99   |
carotene and other perhydro-carotenoids. This is because in the hypersaline anoxic marine and lacustrine environments, the double bond of β-carotene reacts with sulfur and becomes a component of the sulfur cross-linked system.32

The biomarker results show that a low abundance of diasteranes, a high content of gammacerane, and a high homohopane index are signals for a strongly reducing depositional environment. The presence of sulfurized carotenoids in the S1 species suggests a saline lacustrine depositional setting. This result is also consistent with the high content of β-carotene in the total-ion chromatogram (Figure 3). The correlation variance $R^2$ of relative abundance between C$_{30}$ sulfurized carotenoids with DBE$_3$ in S1 compounds and conventional ratios of diasterane/regular sterane and gamma-cerane/C$_{30}$ hopane are 0.98 and 0.99, respectively (Figure 8a,b, Table 1). The excellent relationships between the S1-derived parameters and conventional ratios indicate the application potential of sulfurized carotenoids in sedimentary environmental indicators.

Figure 6. Plots of carbon number vs DBE for the S$_1$ species (a–e) and relative distribution of the S$_1$ species with varying DBE values of the selected oil sands (f–j).

Figure 7. Carbon number distribution and abundance of S$_1$ compounds with DBE values of 1, 3, 6, and 9.
3.2.3. N₁ Class Species and Its Geochemical Significance.

The carbon number and DBE distributions of the N₁ species in the analyzed samples are shown in Figure 9. The carbon number of the N₁ species is distributed in the range of 10−45, mainly distributed between 18 and 35 (Figure 9). The DBE value distribution ranges from 4 to 24. Samples with lower maturity (A₁−A₃) show the DBE distribution mainly in the range 4−14, with DBE values of 7 and 8 as the main peaks (Figure 9a−c,f−h). The DBE value of B₁−B₂ with higher maturity has a wider distribution range within 4−24, with DBE values of 9, 10, and 12 as the main peaks (Figure 9d,e,i−f). As the maturity increases, the carbon number distribution range becomes wider with both lower and higher carbon number compounds, but more compounds are enriched in the DBE value area with a higher condensation degree (Figure 9). The carbon number and DBE distribution of N₁ also reflect the overall impact of the maturity effect.

The carbon number distributions of the main species, the DBE values of 8, 9, 10, and 12, of N₁ compounds are shown in Figure 10. The carbon number is mainly distributed in the range of 10−45, with C₂₈ as the main peak (Figure 10). However, the distribution of the B₂ sample is quite different from that of other samples due to its maturity (Figure 10). The A₁ sample with lower maturity has an absolute predominance of C₂₈ in the compound with the DBE value of 8, and the main peaks of other species are C₂₀ and C₂₈ (Figure 10). The main peaks of carbon of the B₂ sample with higher maturity are C₁₅ and C₁₆ followed by the secondary main peak of C₂₀ (Figure 10), which shows that the N₁ species was affected by maturity. As the maturity increases, the carbon chain is broken, and the main peak of the carbon number moves forward, but the aromatization intensifies (Figure 10).

The N₁ compounds detected by positive-ion ESI were alkylated pyridines and their benzannulated homologues. DBE₇ and DBE₁₀ are likely quinolines and benzoquinolines among other pseudohomologues, respectively, according to the positive-ion ESI mode at low maturity. At the moderate level of maturity, benzoquinolines and pseudohomologues (DBE₁₀) with a more fully aromatic core structure are presented and become predominant within the heteroatom class, as shown in Figures 9e,h and 10d.

The distributions of the pyridine compounds (N₁) of the analyzed oils are quite different. The distribution difference is also influenced by the depositional environment. Pyridine derivatives are often part of biomolecules such as alkaloids and nucleotides. The depositional environment affects the aromatization and comprehensiveness of pyridine compounds. N₁ heteroatom class compounds detected by positive-ion ESI are likely alkylated pyridinic species. DBE values of 5, 7, and 8 represent tetrahydroquinolines, quinolines, and tetrahydroacridines, respectively. Most of the DBE values of 9−15 represent benzoquinolines (DBE₁₀) and indenoquinolines.
lines or azopyrenes (DBE$_{12}$) (Figure 9). The study showed that the DBE ratio of DBE$_{4-8}$/DBE$_{9-15}$ N$_1$ is well-correlated to the sedimentary environment indicators of diasterane/regular sterane and gammacerane/C$_{30}$ hopane ratios (Figure 11). The correlation value $R^2$ between N$_1$DBE$_{4-8}$/DBE$_{9-15}$, and diasterane/regular sterane is 0.9744 (Figure 11a), and the correlation value of $R^2$ between gammacerane/C$_{30}$ hopane reaches 0.999 (Figure 11b), reflecting that the depositional environment has a significant impact on nitrogen species such as pyridines, benzoquinolines, and pseudohomologues. This result shows that the N$_1$-derived parameter can be used as potential indicators for paleoenvironmental evaluation.

3.2.4. O$_1$S$_1$ Class Species and Its Geochemical Significance

The distribution of O$_1$S$_1$ is shown in Figure 12. The DBE values of O$_1$S$_1$ in the analyzed samples are distributed between 0 and 21, mainly 1–12, with DBE values of 1, 2, and 5 as the peak species (Figure 12). As the maturity increases, the compounds are enriched in higher DBE values, mainly distributed in the range 7–16 (Figure 12e,j). The carbon number distribution range of O$_1$S$_1$ for the analyzed samples is between 11 and 45 (Figure 12). As the maturity increases, the carbon number distribution range narrows (Figure 12e), indicating the maturity control on the NSO compound. In general, what cannot be ignored is that the maturity has an overall impact on the composition and distribution of NSO compounds.

5. CONCLUSIONS

Combined with the GC–MS technology, positive-ion (ESI) FT-ICR MS was adopted to understand the composition and distribution characteristics of NSO compounds in the oil sands of the Dongpu Depression. The geochemical significance of the NSO compounds was explored with conventional GC–MS analysis. The relative abundance of nC$_{37}$ in n-alkanes, the appearance of β-carotane, and the high content of gammacerane and C$_{35}$ homohopanes with low content of diasteranes in
the biomarker indicate that the organic matter derives from strongly reducing and saline lacustrine sedimentary environments. The types of NSO compounds in the oil sand extracts in the Dongpu Depression are complex. Nine types of heteroatom compounds have been detected by positive-ion ESI FT-ICR MS, including N, N,N,N,N,S,O,O,S,O, O, O,S, S, and S. The main compounds are S and N compounds, followed by O,S compounds. The DBE value of S species is mainly distributed between 3 and 12, and the carbon number is mainly distributed between 18 and 33, which is correlated to the main carbon number distribution of normal alkanes. The DBE value of N compounds is mainly distributed between 4 and 14, and the carbon number is mainly distributed between 15 and 35. Maturity is one main factor affecting the distribution of NSO heteroatoms in the oil sands. As the maturity increases, the relative abundance of N compounds increases. NSO compounds are enriched in the DBE area with higher condensation, and the main peak carbon shifts forward. This is because the maturity increases and the aromatization intensifies, but the carbon chain is broken and reformed. Among the S species, DBE compounds (thiophenes) have relatively high abundance of C compounds, and their relative abundance has an excellent relationship with the sedimentary environment indicators G/C,H and diasterane/regular sterane. Among the N compounds, DBE/DBE, has a good corresponding relationship with the sedimentary environment indexes G/C,H and diasterane/regular sterane. CDBE S and DBE/DBE, N can be used as references for sedimentary environment identification. The research results play an important role in expanding the application of high-resolution MS technology in the field of petroleum exploration.

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**Notes**

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